WEST Search History

Hide Items Restore Clear Cancel

DATE: Friday, June 23, 2006

Hide?	Set Name	Query	Hit Count
	DB=PGPB, USB	PT,USOC,EPAB,JPAB,DWPI; THES=ASSIGNEE; PLUR=	YES; OP=ADJ
	L6	15 not 11	12
	L5	L4 and tetrafluoroethylene	12
	L4	L2 and ethylene	66
	L3	L2 and copper with catalyst	1
	L2	polyfluoroalkyl acrylate	99
	L1	polyfluoroalkylethyl acrylate	2

END OF SEARCH HISTORY

Welcome to STN International! Enter x:x

LOGINID:ssspta1202jxp

PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

```
Welcome to STN International
NEWS
                Web Page URLs for STN Seminar Schedule - N. America
     1
NEWS
                 "Ask CAS" for self-help around the clock
     2
NEWS 3
        JAN 17 Pre-1988 INPI data added to MARPAT
NEWS 4
        FEB 21
                STN AnaVist, Version 1.1, lets you share your STN AnaVist
                visualization results
NEWS 5
        FEB 22
                The IPC thesaurus added to additional patent databases on STN
       FEB 22 Updates in EPFULL; IPC 8 enhancements added
NEWS 6
     7
NEWS
        FEB 27
                New STN AnaVist pricing effective March 1, 2006
NEWS 8 MAR 03
                Updates in PATDPA; addition of IPC 8 data without attributes
NEWS 9 MAR 22 EMBASE is now updated on a daily basis
NEWS 10 APR 03
                New IPC 8 fields and IPC thesaurus added to PATDPAFULL
NEWS 11 APR 03 Bibliographic data updates resume; new IPC 8 fields and IPC
                thesaurus added in PCTFULL
NEWS 12 APR 04
                STN AnaVist $500 visualization usage credit offered
NEWS 13 APR 12
                LINSPEC, learning database for INSPEC, reloaded and enhanced
NEWS 14 APR 12
                Improved structure highlighting in FQHIT and QHIT display
                in MARPAT
NEWS 15
       APR 12
                Derwent World Patents Index to be reloaded and enhanced during
                second quarter; strategies may be affected
NEWS 16 MAY 10
                CA/CAplus enhanced with 1900-1906 U.S. patent records
NEWS 17 MAY 11
                KOREAPAT updates resume
NEWS 18 MAY 19
                Derwent World Patents Index to be reloaded and enhanced
NEWS 19 MAY 30
                IPC 8 Rolled-up Core codes added to CA/CAplus and
                USPATFULL/USPAT2
        MAY 30
NEWS 20
                The F-Term thesaurus is now available in CA/CAplus
NEWS 21 JUN 02
                The first reclassification of IPC codes now complete in
                INPADOC
NEWS EXPRESS
                FEBRUARY 15 CURRENT VERSION FOR WINDOWS IS V8.01a,
                CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0jc(jp),
                AND CURRENT DISCOVER FILE IS DATED 19 DECEMBER 2005.
                V8.0 AND V8.01 USERS CAN OBTAIN THE UPGRADE TO V8.01a AT
                http://download.cas.org/express/v8.0-Discover/
NEWS HOURS
             STN Operating Hours Plus Help Desk Availability
NEWS LOGIN
             Welcome Banner and News Items
NEWS IPC8
             For general information regarding STN implementation of IPC 8
NEWS X25
             X.25 communication option no longer available after June 2006
```

Enter NEWS followed by the item number or name to see news on that specific topic.

All use of STN is subject to the provisions of the STN Customer agreement. Please note that this agreement limits use to scientific research. Use for software development or design or implementation of commercial gateways or other similar uses is prohibited and may result in loss of user privileges and other penalties.

Welcome to STN International! Enter x:x

LOGINID:ssspta1202jxp

PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

```
Web Page URLs for STN Seminar Schedule - N. America
NEWS
     1
                 "Ask CAS" for self-help around the clock
NEWS
     2
NEWS
        JAN 17
     3
                 Pre-1988 INPI data added to MARPAT
NEWS
        FEB 21
                 STN AnaVist, Version 1.1, lets you share your STN AnaVist
                 visualization results
        FEB 22
NEWS 5
                The IPC thesaurus added to additional patent databases on STN
NEWS 6
        FEB 22 Updates in EPFULL; IPC 8 enhancements added
     7
NEWS
        FEB 27
                New STN AnaVist pricing effective March 1, 2006
NEWS 8 MAR 03 Updates in PATDPA; addition of IPC 8 data without attributes
NEWS 9 MAR 22
                EMBASE is now updated on a daily basis
        APR 03
NEWS 10
                New IPC 8 fields and IPC thesaurus added to PATDPAFULL
        APR 03
NEWS 11
                 Bibliographic data updates resume; new IPC 8 fields and IPC
                 thesaurus added in PCTFULL
NEWS 12
        APR 04
                 STN AnaVist $500 visualization usage credit offered
        APR 12
NEWS 13
                 LINSPEC, learning database for INSPEC, reloaded and enhanced
NEWS 14 APR 12
                 Improved structure highlighting in FQHIT and QHIT display
                 in MARPAT
NEWS 15
        APR 12
                 Derwent World Patents Index to be reloaded and enhanced during
                 second quarter; strategies may be affected
NEWS 16
                 CA/CAplus enhanced with 1900-1906 U.S. patent records
        MAY 10
NEWS 17
        MAY 11
                 KOREAPAT updates resume
NEWS 18 MAY 19
                 Derwent World Patents Index to be reloaded and enhanced
                 IPC 8 Rolled-up Core codes added to CA/CAplus and
NEWS 19 MAY 30
                 USPATFULL/USPAT2
NEWS 20
        MAY 30
                 The F-Term thesaurus is now available in CA/CAplus
NEWS 21
        JUN 02
                 The first reclassification of IPC codes now complete in
                 INPADOC
NEWS EXPRESS
                 FEBRUARY 15 CURRENT VERSION FOR WINDOWS IS V8.01a,
                 CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0jc(jp),
                 AND CURRENT DISCOVER FILE IS DATED 19 DECEMBER 2005.
                 V8.0 AND V8.01 USERS CAN OBTAIN THE UPGRADE TO V8.01a AT
                 http://download.cas.org/express/v8.0-Discover/
NEWS HOURS
              STN Operating Hours Plus Help Desk Availability
NEWS LOGIN
              Welcome Banner and News Items
NEWS IPC8
              For general information regarding STN implementation of IPC 8
NEWS X25
              X.25 communication option no longer available after June 2006
```

Enter NEWS followed by the item number or name to see news on that specific topic.

All use of STN is subject to the provisions of the STN Customer agreement. Please note that this agreement limits use to scientific research. Use for software development or design or implementation of commercial gateways or other similar uses is prohibited and may result in loss of user privileges and other penalties.

: * * * * * * * * * * * * * * * STN Columbus * * * * * * * * * * * * * * * * * *

FILE 'HOME' ENTERED AT 17:54:16 ON 23 JUN 2006

=> file casreact
COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

FILE 'CASREACT' ENTERED AT 17:54:27 ON 23 JUN 2006 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications.

FILE CONTENT:1840 - 18 Jun 2006 VOL 144 ISS 25

New CAS Information Use Policies, enter HELP USAGETERMS for details.

Some CASREACT records are derived from the ZIC/VINITI database (1974-1991) provided by InfoChem, INPI data prior to 1986, and Biotransformations database compiled under the direction of Professor Dr. Klaus Kieslich.

This file contains CAS Registry Numbers for easy and accurate substance identification.

Uploading C:\Program Files\Stnexp\Queries\10516940a.str

L1 STRUCTURE UPLOADED

=> d l1

=>

L1 HAS NO ANSWERS

L1 STR

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

Structure attributes must be viewed using STN Express query preparation.

=> s l1

SAMPLE SEARCH INITIATED 17:55:04 FILE 'CASREACT'
SCREENING COMPLETE - 246 REACTIONS TO VERIFY FROM

25 DOCUMENTS

100.0% DONE 246 VERIFIED 0 HIT RXNS 0 DOCS SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**

PROJECTED VERIFICATIONS: 3980 TO 5860 PROJECTED ANSWERS: 0 TO 0

L2 0 SEA SSS SAM L1 (0 REACTIONS)

=> s l1 full

FULL SEARCH INITIATED 17:55:17 FILE 'CASREACT'

Welcome to STN International! Enter x:x

LOGINID:ssspta1202jxp

PASSWORD:

NEWS X25

TERMINAL (ENTER 1, 2, 3, OR ?):2

```
Welcome to STN International
NEWS
                 Web Page URLs for STN Seminar Schedule - N. America
      1
NEWS
      2
                 "Ask CAS" for self-help around the clock
         JAN 17
                 Pre-1988 INPI data added to MARPAT
NEWS
     3
NEWS
         FEB 21
                 STN AnaVist, Version 1.1, lets you share your STN AnaVist
                 visualization results
NEWS 5
        FEB 22
                 The IPC thesaurus added to additional patent databases on STN
        FEB 22
NEWS
     6
                 Updates in EPFULL; IPC 8 enhancements added
NEWS
     7
         FEB 27
                New STN AnaVist pricing effective March 1, 2006
NEWS
         MAR 03
     8
                Updates in PATDPA; addition of IPC 8 data without attributes
        MAR 22
NEWS 9
                 EMBASE is now updated on a daily basis
NEWS 10
        APR 03
                New IPC 8 fields and IPC thesaurus added to PATDPAFULL
NEWS 11
        APR 03
                Bibliographic data updates resume; new IPC 8 fields and IPC
                 thesaurus added in PCTFULL
NEWS 12
        APR 04
                 STN AnaVist $500 visualization usage credit offered
NEWS 13
        APR 12
                 LINSPEC, learning database for INSPEC, reloaded and enhanced
NEWS 14 APR 12
                 Improved structure highlighting in FQHIT and QHIT display
                 in MARPAT
NEWS 15 APR 12 Derwent World Patents Index to be reloaded and enhanced during
                 second quarter; strategies may be affected
NEWS 16 MAY 10
                 CA/CAplus enhanced with 1900-1906 U.S. patent records
NEWS 17 MAY 11
                KOREAPAT updates resume
NEWS 18 MAY 19
                Derwent World Patents Index to be reloaded and enhanced
NEWS 19 MAY 30
                IPC 8 Rolled-up Core codes added to CA/CAplus and
                 USPATFULL/USPAT2
NEWS 20 MAY 30
                The F-Term thesaurus is now available in CA/CAplus
NEWS 21 JUN 02
                The first reclassification of IPC codes now complete in
                 INPADOC
NEWS EXPRESS
                 FEBRUARY 15 CURRENT VERSION FOR WINDOWS IS V8.01a,
                 CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
                 AND CURRENT DISCOVER FILE IS DATED 19 DECEMBER 2005.
                 V8.0 AND V8.01 USERS CAN OBTAIN THE UPGRADE TO V8.01a AT
                 http://download.cas.org/express/v8.0-Discover/
NEWS HOURS
             STN Operating Hours Plus Help Desk Availability
NEWS LOGIN
             Welcome Banner and News Items
NEWS IPC8
             For general information regarding STN implementation of IPC 8
```

Enter NEWS followed by the item number or name to see news on that specific topic.

All use of STN is subject to the provisions of the STN Customer agreement. Please note that this agreement limits use to scientific research. Use for software development or design or implementation of commercial gateways or other similar uses is prohibited and may result in loss of user privileges and other penalties.

X.25 communication option no longer available after June 2006

* STN Columbus

FILE 'HOME' ENTERED AT 10:32:28 ON 23 JUN 2006

=> file caplus

SINCE FILE COST IN U.S. DOLLARS TOTAL

ENTRY SESSION FULL ESTIMATED COST 0.21 0.21

FILE 'CAPLUS' ENTERED AT 10:32:46 ON 23 JUN 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 23 Jun 2006 VOL 145 ISS 1 FILE LAST UPDATED: 22 Jun 2006 (20060622/ED)

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

http://www.cas.org/infopolicy.html

=> polyfluoroalkylethyl iodide POLYFLUOROALKYLETHYL IS NOT A RECOGNIZED COMMAND The previous command name entered was not recognized by the system. For a list of commands available to you in the current file, enter "HELP COMMANDS" at an arrow prompt (=>).

=> s polyfluoroalkylethyl iodide

1 POLYFLUOROALKYLETHYL

173365 IODIDE

24137 IODIDES

183154 IODIDE

(IODIDE OR IODIDES)

L11 POLYFLUOROALKYLETHYL IODIDE

(POLYFLUOROALKYLETHYL (W) IODIDE)

=> d l1 ibib ab

L1ANSWER 1 OF 1 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2003:1006848 CAPLUS

DOCUMENT NUMBER:

140:17105

TITLE:

Metallic copper catalyst for polyfluoroalkylethyl iodide

production and process for producing

polyfluoroalkylethyl iodides Funakoshi, Yoshirou; Miki, Jun

INVENTOR(S): PATENT ASSIGNEE(S):

Daikin Industries, Ltd., Japan

SOURCE:

PCT Int. Appl., 21 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent Japanese

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

```
DATE
     PATENT NO.
                        KIND
                                         APPLICATION NO.
                               -----
                                           -----
                                        WO 2003-JP7643 20030617
     WO 2003106023
                        A1
                              20031224
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG,
        FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     AU 2003241701
                       A1
                               20031231 AU 2003-241701
                                                               20030617
     CN 1662302
                         Α
                               20050831
                                          CN 2003-814144
                                                                 20030617
     US 2005250966
                        A1
                               20051110
                                          US 2004-516940
                                                                 20041215
PRIORITY APPLN. INFO.:
                                           JP 2002-175381
                                                             A 20020617
                                           WO 2003-JP7643
                                                             W 20030617
OTHER SOURCE(S):
                        MARPAT 140:17105
     Copper catalyzes the addition of ethylene to polyfluoroalkyl iodides to prepare
     polyfluoroalkylethyl iodides. Thus, 138.6 g C2F5I, 8.21
     g Cu, and 1.0 MPa C2H4 were heated 80 min at 80° to give
     perfluoroethylethyl iodide at selectivity 99.8% and C2F5I conversion
     99.5%.
REFERENCE COUNT:
                              THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS
                        32
                              RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
=> s polyfluoroalkyl iodide
          1572 POLYFLUOROALKYL
            1 POLYFLUOROALKYLS
          1572 POLYFLUOROALKYL
                (POLYFLUOROALKYL OR POLYFLUOROALKYLS)
       173365 IODIDE
        24137 IODIDES
       183154 IODIDE
                (IODIDE OR IODIDES)
           89 POLYFLUOROALKYL IODIDE
L2
                (POLYFLUOROALKYL (W) IODIDE)
=> s 12 and ethylene
       524665 ETHYLENE
         3356 ETHYLENES
       526141 ETHYLENE
                (ETHYLENE OR ETHYLENES)
L3
            5 L2 AND ETHYLENE
=> s 13 ibib ab 1-5
MISSING OPERATOR L3 IBIB
The search profile that was entered contains terms or
nested terms that are not separated by a logical operator.
=> d 13 ibib ab 1-5
    ANSWER 1 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
                        2003:1006848 CAPLUS
DOCUMENT NUMBER:
                        140:17105
TITLE:
                        Metallic copper catalyst for polyfluoroalkylethyl
                        iodide production and process for producing
                        polyfluoroalkylethyl iodides
INVENTOR(S):
                        Funakoshi, Yoshirou; Miki, Jun
                       Daikin Industries, Ltd., Japan
PATENT ASSIGNEE(S):
```

SOURCE:

PCT Int. Appl., 21 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT	NO.			KIN	D	DATE		i	APPL	ICAT	ION I	NO.		D	ATE	
					-									-		-
WO 200	31060	23		A1		2003	1224	1	WO 2	003-	JP76	43		2	0030	617
₩:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KR,	ΚZ,	LC,	LK,	LR,	LS,
	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NI,	NO,	NZ,	OM,	PG,
	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	ТJ,	TM,	TN,	TR,	TT,
						VC,										
RW	: GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,
						TM,										
						ΙE,										
	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG
AU 200	32417	01		A1		2003	1231	1	AU 2	003-2	2417	01		2	0030	517
CN 166						2005									0030	517
US 200	52509	66		A1		2005	1110	τ	JS 2	004-5	51694	40		2	0041	215
PRIORITY AP	PLN.	INFO	. :					Ċ	JP 2	002-3	1753	31	I	A 2	0020	517
								V	NO 2	003-0	JP764	43	V	√ 2	0030	517

OTHER SOURCE(S):

MARPAT 140:17105

AB Copper catalyzes the addition of ethylene to polyfluoroalkyl iodides to prepare polyfluoroalkylethyl

iodides. Thus, 138.6 g C2F5I, 8.21 g Cu, and 1.0 MPa C2H4 were heated 80 min at 80° to give perfluoroethylethyl iodide at selectivity 99.8% and C2F5I conversion 99.5%.

REFERENCE COUNT:

THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN

32

ACCESSION NUMBER:

1974:571361 CAPLUS

DOCUMENT NUMBER:

81:171361

TITLE:

SOURCE:

N-(Fluoroalkyl)arylamine dye intermediates

INVENTOR (S):

Sureau, Robert; Pechmeze, Jacques

PATENT ASSIGNEE(S):

Ugine Kuhlmann Ger. Offen., 15 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
DE 2353293	A1	19740516	DE 1973-2353293	19731024	
FR 2205894	A5	19740531	FR 1972-38896	19721103	
CH 590204	A	19770729	CH 1973-14842	19731022	
BE 806852	A1	19740430	BE 1973-137359	19731031	
NL 7314986	A	19740507	NL 1973-14986	19731101	
JP 49108028	A2	19741014	JP 1973-123920	19731102	
GB 1448701	A	19760908	GB 1973-50911	19731102	
IT 999712	Α	19760310	IT 1973-70216	19731105	
PRIORITY APPLN. INFO.:			FR 1972-38896	A 19721103	
AB Twelve RR1N(CH2)2	(CF2) nCF3	(I; R = Ph,	3-MeC6H4, 3-ClC6H4,	or 1-C10H7:	R1
= H, Et, CH2CH2CN,	or CH2C	H2OH; n = 1,	3, or 5) were prepar	ced Thus.	
reaction of PhNH2	with CF3	CF2CH2CH2I ga	ave 80.7 % N- $(3,3,4,4)$	4 -	
pentafluorobutyl)a	aniline (II) [52671-69	5-5]. Similarly prep	pared were 4	other
I. Ethylation of	II gave	84.7% N-ethy	L-N-(3.3.4.4.4-		
pentafluorobutyl)a	niline [52298-66-5].	Similarly prepared	was 1 other	I.

Reaction of II with CH2: CHCN gave 70.2% N-(cyanoethyl)-N-(3,3,4,4,4pentafluorobutyl)aniline [52298-65-4]. Similarly prepared were 2 other I. Reaction of N-(3,3,4,4,4-pentafluorobutyl)-m-toluidine with ethylene oxide gave 75.5% N-(pentafluorobutyl)-N-(hydroxyethyl)-mtoluidine [52671-66-6]. Similarly prepared was 1 other I.

ANSWER 3 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN L3

ACCESSION NUMBER: 1974:519901 CAPLUS

DOCUMENT NUMBER: 81:119901

TITLE: Polyfluoroalkyl iodides

INVENTOR(S): Jaeger, Horst PATENT ASSIGNEE(S): Ciba-Geigy A.-G. Ger. Offen., 36 pp. SOURCE:

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
DE 2361807	A1	19740627	DE 1973-2361807		19731212
CH 576922	Α	19760630	CH 1972-18662		19721221
CA 1023768	A1	19780103	CA 1973-187808		19731210
US 3979469	Α	19760907	US 1973-425011		19731214
GB 1411200	Α	19751022	GB 1973-58328		19731217
FR 2211430	A1	19740719	FR 1973-45565		19731219
BE 808878	A1	19740620	BE 1973-139076		19731220
SU 528029	D	19760905	SU 1973-1978254		19731220
JP 49088811	A2	19740824	JP 1973-142552		19731221
IT 1000773	Α	19760410	IT 1973-54542		19731221
PRIORITY APPLN. INFO.:			CH 1972-18662	A	19721221
			CH 1973-2653	A	19730223
			CH 1973-16037	Α	19731114

AB Twenty-four F3C(CF2)n(CH2CR1F)mCH2CHRI (n = 3-11, m = 1-3, R = H, Me, or CH2OH; R1 = H or F) were prepared by addition reaction of CH2:CHR to F3C(CF2)n(CH2CR1F)mI at 100-60° and 1-30 kg/cm2 in the presence of Al203-supported catalysts containing CuCl, (di)ethanolamine, and optionally SnCl4. Thus, reaction of F3C(CF2)3CH2CF2I with CH2:CH2 over a CuCl-SnCl4-HOCH2CH2NH2 catalyst 7 hr at 140° and 26 kg/cm2 gave 98.5% F3C(CF2)3CH2CF2CH2CH2I.

ANSWER 4 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1971:22415 CAPLUS

DOCUMENT NUMBER:

74:22415

TITLE:

Thermal addition of polyfluoroalkyl

iodides to ethylene

INVENTOR (S):

Bloechl, Walter

PATENT ASSIGNEE(S): SOURCE:

FMC Corp. U.S., 3 pp.

CODEN: USXXAM

Patent

DOCUMENT TYPE:

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
				-	
US 3535393	Α	19701020	US 1968-719692		19680408
PRIORITY APPLN. INFO.:			US 1968-719692	Α	19680408

AB CH2:CH2 (14 g) and 87 g n-C7F15I at 124 ml/min and atmospheric pressure were passed through a Monel spiral preheater at 180-200° and through a Pyrex reactor 60 + 1.5 cm. heated to 379-401° with a spirally-wound Nichrome wire (resistance 2.4 ohms/ft.), residence time 17

sec. The exit gas was condensed to give 91 g n-C7F15CH2CH2I, b20 89-90°, purity 90%. Similarly was prepared 87% n-C8F17CH2CH2I, m. 50°, b10 90-1°.

L3 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1964:410993 CAPLUS

DOCUMENT NUMBER: 61:10993

ORIGINAL REFERENCE NO.: 61:1755d-h,1756a-b

TITLE: Improvement in the preparation of perfluoroalkyl

iodides from tetrafluoroethylene

INVENTOR(S): Parsons, Raymond E.

PATENT ASSIGNEE(S): E. I. du Pont de Nemours & Co.

SOURCE: 5 pp.

DOCUMENT TYPE: Patent
LANGUAGE: Unavailable

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
US 3132185		19640505	US 1962-191722	19620502	
BE 640970 GB 998235			BE GB		
GD 990233			GD		

AB The preparation is given of perfluoroethyl iodide (C2F5I) (I) and longer chain title compds. C2F5(CF2CF2)nI (II), where n = 1 or greater, according to the reaction 2I2 + IF5 + 5CF2:CF2 → 5CF3CF2I. If the molar ratio of iodine to IF5 <2:1, products of formula II (mixture, various n values) result, usually in conjunction with varying amts. of I depending on the iodine to IF5 ratio. Thus, 0.66 mole iodine, 0.33 mole IF5 (III), and 0.0028 mole SbF5 were mixed at ambient temperature in a Hastelloy-C lined autoclave (2 moles iodine and 8.48 millimoles catalyst/mole III), the whole cooled <0°, evacuated to remove air, heated to 80° with agitation and CF2CF2 (IV) and 192 parts added slowly; the pressure rose to 200-50 lb./in.2 gage in <3 min., the temperature rose 8-26° after each addition in about 1 min. and then reverted to about 80° until 165 parts had been added. No temperature increase occurred during the addition of

the

remaining 27 parts of IV. The mixture was kept 1 hr. at 80°, cooled and the volatile products collected in a refrigerated receiver at -60°. Mass-sectrometry, showed 98.3% I, 0.4% unreacted IV, 0.8% IF2CCF2I (V) and 0.4% perfluorocyclobutane (VI). The noncondensed gases contained 28.1 parts IV and 4.3 parts I. The yield of I was 98.5%, based on IV. When the reaction temperature was 60° rather than 80°, the condensed product contained 97.4% I, 0.7% IV, 0.6% VI, 0.7% V, and 0.4% other products; the yield of I was 98.1%. When the reaction was repeated with omission of the SbF5, the condensed product contained 98.5% I, 0.4% IV, 0.5% V and 0.5% VI, the oily residue contained 94.6% V and 4.4% I and the yield of I was 98.4% based on IV. The reaction was repeated with 0.0128 mole anhydrous SnF2; the condensed product contained 96.9% I, 0.6% IV, 0.8% VI, 0.3% V, and 1.3% other products. The yield of I was 99% based on IV. Similarly, other catalysts were used (catalyst, and amount (parts) and % yield of I given): CaF2, 2.0, 78; CrF3.3H2O, 2.0, 16; KF, 2.0, 18; HF, 100.0, 15; CoF2.2H2O, 2.0, 60; PbF4, 2.0, 39; CuF2, 2.0, 78; NiF2.5H2O, 2.0, 28; and AgF (50%), 2.0, 73. The following examples give the mixture of components. A mixture of 0.33 mole iodine, 0.182 mole III, and 0.0028 mole SbF3 (VII) was prepared at ambient temperature in a Hastelloy-C lined autoclave,

cooled to below 0°, evacuated, sealed, heated to 60° with agitation, and 1.21 moles IV added slowly in small increments in 1 hr. The reaction was exothermic, the pressure dropped rapidly from 150-250 lb./in.2 gage to <100 lb./in.2 gage, the temperature rose 7-44° after each addition The mixture was kept at 70° 1 hr., cooled, the volatile products were vented through a H2O scrubber and a CaSO4 drying column, condensed in a refrigerated receiver at -60° and the condensed volatile material analyzed. It contained (in mole-%) 95.1 I, 1.2 C4F9I

(VIII), 0.5 IV, and 1 C2F6 (IX). The noncondensed gases were found to consist of 8.0 parts IV, 4.0 parts I, and 3.1 parts IX. The nonvolatile residue (100 parts) was washed with H2O and analyzed with a vapor phase chromatograph (in weight-%): I 21.1; VIII 28.7; C6F13I (X) 20.6; C8F17I (XI) 13.0; C10F21I (XII) 7.5; C12F25I (XIII) 4.1; C14F29I (XIV) 2.3; C16F33I (XV) 1.1. The combination of these analyses indicates that 96.9% IV is accounted for; % yields are as follows (based on IV consumed): I 48.8, VIII 15.8, X 12.3, XI 1.7, XV 0.9, and IX 0.6. Thus, the total yield of I was 95.8% and the total yield of products II was 47.1%. Iodine consumption was quant. Similarly, addnl. expts. were carried out with a varying ratio of iodine and III, as well as the temperature and the catalysts.

=> d his

(FILE 'HOME' ENTERED AT 10:32:28 ON 23 JUN 2006)

FILE 'CAPLUS' ENTERED AT 10:32:46 ON 23 JUN 2006

L1 1 S POLYFLUOROALKYLETHYL IODIDE

L2 89 S POLYFLUOROALKYL IODIDE

L3 5 S L2 AND ETHYLENE

=> s tetrafluoroethylene and 12

25843 TETRAFLUOROETHYLENE

34 TETRAFLUOROETHYLENES

25850 TETRAFLUOROETHYLENE

(TETRAFLUOROETHYLENE OR TETRAFLUOROETHYLENES)

L4 8 TETRAFLUOROETHYLENE AND L2

=> s 14 not 13

L5 6 L4 NOT L3

=> d 15 ibib ab 1-6

L5 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:381429 CAPLUS

DOCUMENT NUMBER: 127:81136

TITLE: Pentafluoro-λ6-sulfanyl (SF5) fluoroalkyl

iodides

AUTHOR(S): Terjeson, Robin J.; Renn, Julia; Winter, Rolf; Gard,

Gary L.

CORPORATE SOURCE: Dep. Chem., Portland State Univ., Portland, OR, 97207,

USA

SOURCE: Journal of Fluorine Chemistry (1997), 82(1), 73-78

CODEN: JFLCAR; ISSN: 0022-1139

PUBLISHER: Elsevier
DOCUMENT TYPE: Journal
LANGUAGE: English

OTHER SOURCE(S): CASREACT 127:81136

AB The synthesis of several SF5-containing fluoroalkyl iodides derived from an improved synthesis of SF5CF2CF2I are reported. These include

SF5CF2CF2CH2CH2I, SF5(CF2)4I, SF5CF2CF2(CH2)4I, SF5CF2CF2CH:CHI and

SF5CF2CF2CHFCF21. The higher homologs SF5(CF2)61, SF5CF2CF2(CHFCF2)21 and

SF5CF2CF2(CH2CH2)3I were also identified.

REFERENCE COUNT: 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 2 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:137648 CAPLUS

DOCUMENT NUMBER: 124:177222

TITLE: Initiators for telomerization of

polyfluoroalkyl iodides with

fluoroolefins

INVENTOR(S): Krespan, Carl George; Petrov, Viacheslav Alexandrovic;

Smart, Bruce Edmund

חאתה

PATENT ASSIGNEE(S): du Pont de Nemours, E. I., and Co., USA

SOURCE: PCT Int. Appl., 14 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

AUTHOR(S):

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9532936	A1	19951207	WO 1995-US5857	19950518
W: AU, CA, CN	JP, KR			
RW: AT, BE, CH	DE, DK	, ES, FR,	GB, GR, IE, IT, LU, MC	, NL, PT, SE
US 5574193	A	19961112	US 1995-416942	19950404
AU 9525857	A1	19951221	AU 1995-25857	19950518
PRIORITY APPLN. INFO.:			US 1994-249311	A 19940526
			WO 1995-US5857	W 19950518

AB Polyfluoroalkyl iodides are telomerized with polyfluoroolefins in the presence of strong fluorooxidizer initiators. The products obtained are useful in the production of surfactants and oil repellents. Tetrafluoroethylene and perfluorobutyl iodide were telomerized in the presence of XeF2.

L5 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

KIMD

ACCESSION NUMBER: 1988:21312 CAPLUS

DOCUMENT NUMBER: 108:21312

TITLE: Studies of fluoroalkylation and fluoroalkoxylation.

Part 16. Reactions of fluoroalkyl iodides with some

ADDITONTION NO

חאייינים

nucleophiles by SRN1 mechanism Chen, Qing Yun; Qiu, Zai Ming

CORPORATE SOURCE: Shanghai Inst. Org. Chem., Acad. Sin., Shanghai, Peop.

Rep. China

SOURCE: Journal of Fluorine Chemistry (1987), 35(2), 343-57

CODEN: JFLCAR; ISSN: 0022-1139

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 108:21312

AB XCF2CF2I [I; X = ClCF2CF2, Cl(CF2)4] readily react with the anion of Et acetoacetate in DMF to give XCF2C(CH2CO2Et):CHCO2Et (II), XCF2CF2H (III), Ac2CHCO2Et, and AcCH(CO2Et)2CH(CO2Et)Ac (IV). The reaction can be partly suppressed with 1,4-dinitrobenzene and the radical intermediate can be trapped by diallyl ether (DAE) to give the THF derivs. Anions of acetylacetone and malonitrile react also with I in the presence of DAE to yield the five-membered ring compds. All these results seem to indicate that the reaction is a radical chain process induced by electron transfer. In the case of XCF2CF2I (I; X = Cl), tetrafluoroethylene and IV, instead of II and III, are produced, showing the occurrence of β-fragmentation of the 2-halotetrafluoroethyl radical in the initiation step.

L5 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1977:422253 CAPLUS

DOCUMENT NUMBER: 87:22253

TITLE: Methyl-terminated perfluoroalkyl iodides and related

compounds

AUTHOR(S): Rondestvedt, Christian S., Jr.

CORPORATE SOURCE: Org. Chem. Dep., E. I. du Pont de Nemours and Co.,

Wilmington, DE, USA

SOURCE: Journal of Organic Chemistry (1977), 42(11), 1985-90

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: English

AB 1,1-Difluoroethyl iodide was prepared from vinylidene fluoride and hydrogen

iodide in 96% yield; it telomerized with tetrafluoroethylene with acyl peroxide catalysis to give the resp. Me(CF2)nI (n = 3,5,7,9,11,13,15).

ANSWER 5 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1970:476644 CAPLUS

DOCUMENT NUMBER:

73:76644

TITLE:

Linear polyfluoroalkyl iodide

compounds

INVENTOR(S):

Rebsdat, Siegfried; Schuierer, Erich; Hahn, Helmut

PATENT ASSIGNEE(S): Farbwerke Hoechst A.-G.

SOURCE:

Ger., 4 pp. CODEN: GWXXAW

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 1915395	Α	19700702	DE 1969-1915395	19690326
NL 7004025	Α	19700929	NL 1970-4025	19700320
BE 748010	Α	19700928	BE 1970-748010	19700326
FR 2035913	A5	19701224	FR 1970-10881	19700326
FR 2035913	B1	19740503		
GB 1256818	Α	19711215	GB 1970-1256818	19700326
PRIORITY APPLN. INFO.:			DE 1969-1915395 A	19690326
AB Compds. of the gene	ral for	cmula, C2F5(CF2CF2)nI, where $n = 2$	to 5, were
prepared by treating	g C2F5]	[with tetra:	fluoroethylene in the	
presence of 0.08-0.	2% bis	(trichloroac:	ryloyl) peroxide, at 60	-75°
			e intermediates for the	
hydro-		_		•

and oleophobic media.

ANSWER 6 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1969:11105 CAPLUS

DOCUMENT NUMBER:

70:11105

TITLE:

Polyfluoroalkyl iodides from tetrafluoroethylene and lower

polyfluoroalkyl iodides

INVENTOR(S):

Blochl, Walter

PATENT ASSIGNEE(S): SOURCE:

FMC Corp. U.S., 5 pp. CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 3404189	Α	19681001	US 1966-603072	19661104
PRIORITY APPLN. INFO.:			US 1966-603072 A	19661104
AB A 3:1 molar mixture	of 29.	1 g. n-C7H15	I (I) and Fe2C:CF2 were	heated to
200° and passed thr	ough a	glass tube 1	.5 m. + 6 mm. at	
355° and 80 mm., re	sidence	time 12 min	., to give a condensate	of
26.65 g. I, 3.64 g.	n-C9F1	9I, and 0.53	g. 90% n-C11F23I. Sim	ilarly were
prepared 13% H(CF2)	4I, b75	0 85°; 86% C	l(CF2)4I, b350 70°; 2.7	•
g. I(CF2)4I, from 2	0 g. IC	F2CF2I, b100	86°; 72% Br(CF2)4I. b.	
130°; and a mixture	of 80%	n-C10F21I a	nd 15% n-C12F25I. A dr	awing of
the apparatus is in	cluded.			J

(FILE 'HOME' ENTERED AT 10:32:28 ON 23 JUN 2006)

FILE 'CAPLUS' ENTERED AT 10:32:46 ON 23 JUN 2006 L11 S POLYFLUOROALKYLETHYL IODIDE 89 S POLYFLUOROALKYL IODIDE L2L3 5 S L2 AND ETHYLENE 8 S TETRAFLUOROETHYLENE AND L2 **L4** L5 6 S L4 NOT L3 => s polyfluoroalkyl acrylate 1572 POLYFLUOROALKYL 1 POLYFLUOROALKYLS 1572 POLYFLUOROALKYL (POLYFLUOROALKYL OR POLYFLUOROALKYLS) 180045 ACRYLATE 35260 ACRYLATES 189689 ACRYLATE (ACRYLATE OR ACRYLATES) L672 POLYFLUOROALKYL ACRYLATE (POLYFLUOROALKYL (W) ACRYLATE) => s 16 and tetrafluoroethylene 25843 TETRAFLUOROETHYLENE 34 TETRAFLUOROETHYLENES 25850 TETRAFLUOROETHYLENE (TETRAFLUOROETHYLENE OR TETRAFLUOROETHYLENES) L7 1 L6 AND TETRAFLUOROETHYLENE => d 17 ibib ab ANSWER 1 OF 1 CAPLUS COPYRIGHT 2006 ACS on STN L7ACCESSION NUMBER: 1993:214334 CAPLUS DOCUMENT NUMBER: 118:214334 TITLE: UV absorbers based on fluorine-containing acrylic polymers INVENTOR(S): Kodama, Shunichi; Kawasaki, Toru; Kobayashi, Shigeki; Funaki, Atsushi Asahi Glass Co., Ltd., Japan PATENT ASSIGNEE(S): SOURCE: Jpn. Kokai Tokkyo Koho, 5 pp. CODEN: JKXXAF DOCUMENT TYPE: Patent Japanese LANGUAGE: FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE ---------_____ -----_____ JP 04288387 A2 19921013 JP 1991-75681 19910315 JP 2980999 B2 19991122 PRIORITY APPLN. INFO.: JP 1991-75681 Title polymers are obtained from radical-polymerizable double bond-containing compds. [for example, (meth)acryloyl group-containing benzophenones, Ph salicylates, or benzotriazoles] and C4-21-polyfluoroalkyl or polyfluoroether-containing (meth)acrylates at mol ratio 90/10 - 10/90 and show weight average mol. weight (Mw) 1000-100,000. Thus, autoclaving 71 parts 2-(3'-hydroxy-4'-benzoyl)phenyloxyethyl methacrylate (I) and 126 parts perfluoroalkylethyl acrylate (II) in Me2CO in the presence of peroxybutyl isobutyrate under N at 65° gave a copolymer (III) with Mw .apprx.5000. A 25 μm-thick film extruded from a mixture of 100 parts ethylene-chlorotrifluoroethylene-tetrafluoroethylene copolymer and 5 parts of the III showed UV permeation 0% at 340 nm and 0.2% at 360

nm initially and 0 and 0.3%, resp., after 1000-h exposure to a Sunshine

arc weatherometer.

WEST Search History

Hide Items | Restore | Clear | Cancel |

DATE: Friday, June 23, 2006

Hide?	Set Name	Query	<u>Hit</u>
	<u>Name</u>		Count
	DB=PC	GPB, USPT, USOC, EPAB, JPAB, DWPI; THES=ASSIGNEE; PLUR=YES; OP=A	1DJ
	L13	L12 or 110	68
	L12	L11 and ethylene	47
	L11	method with (produc\$3 or prepar\$3 or mak\$3)same (polyfluoroalkyl iodide or perfluoroalkyl iodide)	110
	L10	L9 not 15 not 16	34
	L9	L8 not 11	34
	L8	L7 and ethylene	35
	L7	process with (produc\$3 or prepar\$3 or mak\$3)same (polyfluoroalkyl iodide or perfluoroalkyl iodide)	109
	L6	15 not 11	12
	L5	L4 and tetrafluoroethylene	12
	L4	L2 and ethylene	66
П.	L3	L2 and copper with catalyst	1
	L2	polyfluoroalkyl acrylate	99
	L1	polyfluoroalkylethyl acrylate	2

END OF SEARCH HISTORY

WEST Search History

Hide liems Restore Clear Cancel

DATE: Friday, June 23, 2006

Hide?	<u>Set</u> Name	Query	<u>Hit</u> Count
	DB=PC	GPB,USPT,USOC,EPAB,JPAB,DWPI; THES=ASSIGNEE; PLUR=YES; OP=	ADJ
	L14	113 and copper	14
	L13	L12 or 110	68
	L12	L11 and ethylene	47
	L11	method with (produc\$3 or prepar\$3 or mak\$3)same (polyfluoroalkyl iodide or perfluoroalkyl iodide)	110
	L10	L9 not 15 not 16	34
	L9	L8 not l1	34
	L8	L7 and ethylene	35
	L7	process with (produc\$3 or prepar\$3 or mak\$3)same (polyfluoroalkyl iodide or perfluoroalkyl iodide)	109
	L6	15 not 11 .	12
	L5	L4 and tetrafluoroethylene	12
	L4	L2 and ethylene	66
	L3	L2 and copper with catalyst	1
	L2	polyfluoroalkyl acrylate	99
	L1	polyfluoroalkylethyl acrylate	2

END OF SEARCH HISTORY

Welcome to STN International! Enter x:x

LOGINID:ssspta1202jxp

PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

```
* * * * * * * * *
                     Welcome to STN International
NEWS 1
                 Web Page URLs for STN Seminar Schedule - N. America
NEWS 2
                 "Ask CAS" for self-help around the clock
NEWS 3 JAN 17
                 Pre-1988 INPI data added to MARPAT
NEWS 4 FEB 21 STN AnaVist, Version 1.1, lets you share your STN AnaVist
                 visualization results
NEWS 5 FEB 22 The IPC thesaurus added to additional patent databases on STN
NEWS 6 FEB 22 Updates in EPFULL; IPC 8 enhancements added
NEWS 7 FEB 27 New STN AnaVist pricing effective March 1, 2006
NEWS 8 MAR 03 Updates in PATDPA; addition of IPC 8 data without attributes
NEWS 9 MAR 22
                 EMBASE is now updated on a daily basis
NEWS 10 APR 03
                 New IPC 8 fields and IPC thesaurus added to PATDPAFULL
NEWS 11 APR 03
                 Bibliographic data updates resume; new IPC 8 fields and IPC
                 thesaurus added in PCTFULL
NEWS 12 APR 04
                 STN AnaVist $500 visualization usage credit offered
NEWS 13 APR 12
                 LINSPEC, learning database for INSPEC, reloaded and enhanced
NEWS 14 APR 12
                 Improved structure highlighting in FQHIT and QHIT display
                 in MARPAT
NEWS 15 APR 12
                 Derwent World Patents Index to be reloaded and enhanced during
                 second quarter; strategies may be affected
NEWS 16 MAY 10
                 CA/CAplus enhanced with 1900-1906 U.S. patent records
NEWS 17 MAY 11
                 KOREAPAT updates resume
NEWS 18 MAY 19
                 Derwent World Patents Index to be reloaded and enhanced
NEWS 19 MAY 30 IPC 8 Rolled-up Core codes added to CA/CAplus and
                 USPATFULL/USPAT2
NEWS 20 MAY 30
                 The F-Term thesaurus is now available in CA/CAplus
NEWS 21
        JUN 02
                 The first reclassification of IPC codes now complete in
                 INPADOC
NEWS EXPRESS
                 FEBRUARY 15 CURRENT VERSION FOR WINDOWS IS V8.01a,
                 CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
                 AND CURRENT DISCOVER FILE IS DATED 19 DECEMBER 2005.
                 V8.0 AND V8.01 USERS CAN OBTAIN THE UPGRADE TO V8.01a AT
                 http://download.cas.org/express/v8.0-Discover/
NEWS HOURS
              STN Operating Hours Plus Help Desk Availability
NEWS LOGIN
              Welcome Banner and News Items
NEWS IPC8
              For general information regarding STN implementation of IPC 8
NEWS X25
              X.25 communication option no longer available after June 2006
```

Enter NEWS followed by the item number or name to see news on that specific topic.

All use of STN is subject to the provisions of the STN Customer agreement. Please note that this agreement limits use to scientific research. Use for software development or design or implementation of commercial gateways or other similar uses is prohibited and may

result in loss of user privileges and other penalties.

FILE 'HOME' ENTERED AT 17:54:16 ON 23 JUN 2006

=> file casreact COST IN U.S. DOLLARS

SINCE FILE TOTAL. ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

FILE 'CASREACT' ENTERED AT 17:54:27 ON 23 JUN 2006 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications.

FILE CONTENT: 1840 - 18 Jun 2006 VOL 144 ISS 25

New CAS Information Use Policies, enter HELP USAGETERMS for details.

***************** CASREACT now has more than 10 million reactions *****************

Some CASREACT records are derived from the ZIC/VINITI database (1974-1991) provided by InfoChem, INPI data prior to 1986, and Biotransformations database compiled under the direction of Professor Dr. Klaus Kieslich.

This file contains CAS Registry Numbers for easy and accurate substance identification.

Uploading C:\Program Files\Stnexp\Queries\10516940a.str

L1STRUCTURE UPLOADED

=> d 11

L1 HAS NO ANSWERS

STR

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 17:55:04 FILE 'CASREACT' SCREENING COMPLETE -246 REACTIONS TO VERIFY FROM 25 DOCUMENTS

246 VERIFIED 100.0% DONE 0 HIT RXNS 0 DOCS SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE**

PROJECTED VERIFICATIONS: 3980 TO 5860 PROJECTED ANSWERS: O TO

L2

=> s 11 full

FULL SEARCH INITIATED 17:55:17 FILE 'CASREACT'

SCREENING COMPLETE - 4448 REACTIONS TO VERIFY FROM 437 DOCUMENTS

100.0% DONE 4448 VERIFIED 0 HIT RXNS 0 DOCS

SEARCH TIME: 00.00.04

L3 0 SEA SSS FUL L1 (0 REACTIONS)

=> s ll sss full

FULL SEARCH INITIATED 17:55:36 FILE 'CASREACT'

SCREENING COMPLETE - 4448 REACTIONS TO VERIFY FROM 437 DOCUMENTS

100.0% DONE 4448 VERIFIED 0 HIT RXNS 0 DOCS

SEARCH TIME: 00.00.04

L4 0 SEA SSS FUL L1 (0 REACTIONS)

=> file caplus

COST IN U.S. DOLLARS SINCE FILE TOTAL

ENTRY SESSION

FULL ESTIMATED COST 220.72 220.93

FILE 'CAPLUS' ENTERED AT 17:56:00 ON 23 JUN 2006
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 23 Jun 2006 VOL 145 ISS 1 FILE LAST UPDATED: 22 Jun 2006 (20060622/ED)

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

http://www.cas.org/infopolicy.html

=> s ethylene and perfluoroalkyl iodide

524665 ETHYLENE

3356 ETHYLENES

526141 ETHYLENE

(ETHYLENE OR ETHYLENES)

10971 PERFLUOROALKYL

35 PERFLUOROALKYLS

10991 PERFLUOROALKYL

(PERFLUOROALKYL OR PERFLUOROALKYLS)

173365 IODIDE

24137 IODIDES

183154 IODIDE

(IODIDE OR IODIDES)

=> s 15 and copper

890360 COPPER

436 COPPERS

890423 COPPER

(COPPER OR COPPERS)

4 L5 AND COPPER L6

=> d 16 ibib ab 1-4

ANSWER 1 OF 4 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:125104 CAPLUS

DOCUMENT NUMBER: 140:320999

TITLE: Environmentally Benign Processes for Making Useful

Fluorocarbons: Nickel- or Copper(I)

Iodide-Catalyzed Reaction of Highly Fluorinated Epoxides with Halogens in the Absence of Solvent and

Thermal Addition of CF2I2 to Olefins

AUTHOR(S): Yang, Zhen-Yu

CORPORATE SOURCE: Experimental Station, Central Research Development, E.

I. Du Pont de Nemours and Co., Wilmington, DE,

19880-0328, USA

SOURCE: Journal of Organic Chemistry (2004), 69(7), 2394-2403

CODEN: JOCEAH; ISSN: 0022-3263

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 140:320999

Highly fluorinated epoxides react with halogens in the presence of nickel powder or CuI at elevated temps. to provide a useful and general synthesis of dihalodifluoromethanes (CF2X2) and fluoroacyl fluorides in the absence of solvent. At 185 °C, hexafluoropropylene oxide and halogens produce CF2X2 (X = I, Br) in 68-90% isolated yields, along with small amts. of X(CF2)nX, (n = 2, 3). With interhalogens I-X (X = C1, Br), a mixture of CF2I2, CF2XI, and CF2X2 was obtained. The fluorinated epoxides substituted with perfluorophenyl, fluorosulfonyl, and chlorofluoroalkyl groups also react cleanly with iodine to give CF2I2 and the corresponding fluorinated acyl fluorides in good yields. The reaction probably involves an oxidative addition of fluorinated epoxides into metal surfaces to form an oxametallacycle, followed by rapid decomposition to difluorocarbene-metal surfaces, which alters the reactivity of the difluorocarbene carbon from electrophilic to nucleophilic. The increase of nucleophilicity of difluorocarbene facilitates the reaction with electrophilic halogens. CF2I2 reacted with olefins thermally to give 1,3-diiodofluoropropane derivs. Both fluorinated and non-fluorinated alkenes gave good yields of the adducts. Reaction with ethylene, propylene, perfluoroalkylethylene, vinylidene fluoride, and trifluoroethylene provided the corresponding adducts in 58-86% yields. With tetrafluoroethylene, a 1:1 adduct was predominantly formed along with small amts. of higher homologs. In contrast to perfluoroalkyl iodides, CF2I2 also readily adds to perfluorovinyl ethers to give 1,3-diiodoperfluoro ethers. The formation of fluorocarbene was discussed.

REFERENCE COUNT: 117 THERE ARE 117 CITED REFERENCES AVAILABLE FOR

THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 2 OF 4 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1984:591104 CAPLUS

DOCUMENT NUMBER: 101:191104

Reactivity of perfluoroiodoalkanes with alkyl TITLE:

carbonates and pyrocarbonates in the presence of a

zinc-copper couple

Benefice, S.; Blancou, H.; Commeyras, A. AUTHOR(S):

Lab. Chim. Org., Univ. Sci. Tech. Languedoc, CORPORATE SOURCE:

Montpellier, 34060, Fr.

Tetrahedron (1984), 40(9), 1541-4 SOURCE:

CODEN: TETRAB; ISSN: 0040-4020

DOCUMENT TYPE: Journal French LANGUAGE:

OTHER SOURCE(S): CASREACT 101:191104

Perfluoroalkyl iodides RI [R = F3C(CF2)n; n = 3, 5, 7]reacted with Zn-Cr in ethylene carbonate at 80-90° to

give 60-80% RCO2CH2CH2OH, and at >150° to give 60-85% RCO2H. RI

reacted with Cu-Zn in (EtO)2CO or (EtO2C)2O to give 50-70, 50-60% RCO2Et,

resp. In all cases the coupling product RR was also formed.

ANSWER 3 OF 4 CAPLUS COPYRIGHT 2006 ACS on STN

1974:81946 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 80:81946

TITLE: Reaction of (perfluoroalkyl)copper compounds

with 1-bromo-1-(perfluoroalkyl)ethylenes

Santini, G.; Le Blanc, M.; Reiss, J. G. AUTHOR(S):

Dep. Chim., Inst. Math. Sci. Phys., Nice, Fr. CORPORATE SOURCE: SOURCE:

Tetrahedron (1973), 29(16), 2411-14

CODEN: TETRAB; ISSN: 0040-4020

DOCUMENT TYPE: Journal LANGUAGE: English

Heating RCBr:CH2 [I, R = (CF2)5CF3, (CF2)7CF3], R1I [R = (CF2)3CF3,

(CF2)5CF3, (CF2)7CF3], precipitated Cu, and DMF for 20 hr at 120-40° in a

sealed tube gave 60-70% trans-RCH:CHR1. I were prepared by

bromination-dehydrobromination of RCH: CH2. 1H and 19F NMR were determined

ANSWER 4 OF 4 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1961:33124 CAPLUS

DOCUMENT NUMBER: 55:33124

ORIGINAL REFERENCE NO.: 55:6496h-i,6497a-c

TITLE: Perfluoroalkylated aromatic compounds

PATENT ASSIGNEE(S): Minnesota Mining and Manufacturing Co.

DOCUMENT TYPE: Patent LANGUAGE: Unavailable

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. _____ GB 840725 19600706 GB 1956-24567 19560810

Aromatic compds. were perfluoroalkylated with the appropriate perfluoroalkyl iodide. Some of the products were converted to dyes. The following reactions were run in a 180 ml. stainless steel rocking autoclave capped by a 3000 lb./sq. in. Ni rupture disk. n-C3F7I (90 g.) and 23.4 g. C6H6 was heated 15 hrs. at 250°, the crude product filtered off, treated with Hg and distilled to yield 22.3 g. C3F7Ph, b760 132°, n25D 1.3790, and 1.5 g. isomeric (C3F7)2C6H4, n25D 1.3492. Likewise prepared were C7F15Ph (32% yield), b760 200 n25D 1.3596, and (C7F15)2C6H4 (4% yield), m. 88°. All subsequent reactions were run in heavy-walled 30 ml. glass ampuls. n-C7F15I (10 g.) and 0.9 g. PhMe was heated 14 hrs. at 290°, the mixture extracted with perfluorooxacycloheptane and distilled to give a 31% yield of C7F15C6H4Me, b760 217°, n25D 1.3678. Similarly prepared were: C7F15C10H7 in 31% yield (from naphthalene), n25D 1.4160; C7F15C6H4CN in 26% yield (from

PhcN), b760 254°, m. 45-52°; C7F15C6H4Br in 30% yield (from PhBr), b760 237°, n25D 1.3870; C7F15C6H4I in 38% yield (from PhI), b760 253°, n25D 1.4152; C7F15C6H3(CO)2O (I) in 36% yield (from C6H4(CO)2O), m. 121-4°. I and glycerol in 3:2 mole ratio heated 4 hrs. at 220° gave fluorinated alkyd resins that had varnishlike properties. A perfluoroalkylated dye was prepared as follows: a 30 ml. glass ampul was charged with 1.4 g. Cu phthalocyanine and 10 g. n-C7F15I. The sealed ampul was heated 14 hrs. at 330°. Extraction with perfluorooxacycloheptane gave 4.4 g. C60F60H12N8Cu (I) (purity approx. 85%). Impurities were removed by vacuum sublimation to yield I, m. 300°, visible absorption at 6550-90 A. and 6200 A. I was soluble in fluorinated solvents only. Similarly perfluoroalkylated were the following: thioindigo (Color Index 1207), pyranthrone (Color Index 1096), violanthrone (Color Index 1099), and dichloroisoviolanthrone (Color Index 1104). The principal visible absorption peaks of the products were given. These dyes have been used to dye Teflon.

=> d his

(FILE 'HOME' ENTERED AT 17:54:16 ON 23 JUN 2006)

FILE 'CASREACT' ENTERED AT 17:54:27 ON 23 JUN 2006

L1 STRUCTURE UPLOADED

L2 0 S L1

L3 0 S L1 FULL

L4 0 S L1 SSS FULL

FILE 'CAPLUS' ENTERED AT 17:56:00 ON 23 JUN 2006

L5 35 S ETHYLENE AND PERFLUOROALKYL IODIDE

L6 4 S L5 AND COPPER

=> s 15 not 16

L7 31 L5 NOT L6

=> d 17 ibib ab 1-31

L7 ANSWER 1 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:735265 CAPLUS

DOCUMENT NUMBER: 143:220367

TITLE: Method of fabricating dual damascene interconnection

and etchant for stripping sacrificial layer

INVENTOR(S): Han, Sang-cheol; Lee, Kyoung-woo; Kim, Mi-young

PATENT ASSIGNEE(S): Samsung Electronics Co., Ltd., S. Korea

SOURCE: U.S. Pat. Appl. Publ., 13 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
US 2005176243	A1	20050811,	US 2005-33208		20050111
PRIORITY APPLN. INFO.:			KR 2004-8065	Α	20040206
OTHER SOURCE(S):	MARPAT	143:220367			

AB A method of forming a dual damascene semiconductor interconnection and an etchant composition specially adapted for stripping a sacrificial layer in a dual damascene fabrication process without profile damage to a dual damascene pattern are provided. The method includes sequentially forming a 1st etch stop layer, a 1st intermetal dielec., a 2nd intermetal dielec., and a capping layer on a surface of a semiconductor substrate on which a

lower metal wiring is formed; etching the 1st intermetal dielec., the 2nd intermetal dielec., and the capping layer to form a via; forming a sacrificial layer within the via; etching the sacrificial layer, the 2nd intermetal dielec., and the capping layer to form a trench; removing the sacrificial layer remaining around the via using an etchant composition including NH4F, HF, H2O and a surfactant; and forming an upper metal wiring within the thus formed dual damascene pattern including the via and the trench. The preferred etchant composition for stripping a sacrificial layer in the foregoing dual damascene process consists essentially of NH4F, HF, H2O and a surfactant.

L7 ANSWER 2 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:727519 CAPLUS

DOCUMENT NUMBER: 144:312852

TITLE: Amphiphilic block copolymers having water soluble and

perfluoroalkyl-group containing blocks

AUTHOR(S): Kressler, Joerg; Kaiser, Sergej

CORPORATE SOURCE: Department of Engineering Science Martin-Luther-

Universitaet Halle-Wittenberg, Halle, D-06099, Germany

SOURCE: Polymer Preprints (American Chemical Society, Division

of Polymer Chemistry) (2005), 46(2), 580-581

CODEN: ACPPAY; ISSN: 0032-3934

PUBLISHER: American Chemical Society, Division of Polymer

Chemistry

DOCUMENT TYPE: Journal; (computer optical disk)

LANGUAGE: English

AB A serious of amphiphilic block copolymer having polyethylene oxide as the

water soluble block were synthesized, and then esterified with

perfluoroalkyl iodide. A second triblock copolymer was

synthesized from ethylene oxide and hexafluoropropylene oxide.

The aggregation behavior of the polymers and the interactions of the above block polymer with lipid monomers were also studied by surface pressure

measurement and IRRAS.

REFERENCE COUNT:

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS

L7 ANSWER 3 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:769016 CAPLUS

DOCUMENT NUMBER: 141:395229

TITLE: An easy three step synthesis of perfluoroalkylated

amphetamines

AUTHOR(S): Tewari, Amit; Hein, Martin; Zapf, Alexander; Beller,

Matthias

CORPORATE SOURCE: Universitaet Rostock, Rostock, 18059, Germany

SOURCE: Tetrahedron Letters (2004), 45(41), 7703-7707

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER: Elsevier B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 141:395229

AB A general synthesis of perfluoroalkylated amphetamines is presented.

Initially, 1-aryl-1-iodo-2-(perfluoroalkyl)ethylenes are prepared

by radical addition of perfluoroalkyl iodides to

arylacetylenes. Key step of the reaction sequence is the following dehydroiodination in the presence of n-BuLi to give 1-perfluoroalkyl-2-arylacetylenes in situ, which are reacted with secondary amines to produce perfluoroalkylated enamines in a new one pot procedure. Final

hydrogenation yields the desired products in good yields. By using N,N-dibenzylamine or N-benzylamines the corresponding primary and

secondary perfluoroalkylated amines are easily available.

REFERENCE COUNT: 44 THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 4 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:385293 CAPLUS

DOCUMENT NUMBER: 141:349790

TITLE: Pentafluoro-λ6-sulfanyl (SF5)

perfluoroalkyl iodides-synthesis and

reaction with ethylene and

tetrafluoroethylene. Crystal structure of

SF5 (CF2) 4CH2CH2I

Nixon, Paul G.; Mohtasham, Javid; Winter, Rolf; Gard, AUTHOR(S):

Gary L.; Twamley, Brendan; Shreeve, Jean'ne. M.

CORPORATE SOURCE: Department of Chemistry, Portland State University,

Portland, OR, 97207-0751, USA

SOURCE: Journal of Fluorine Chemistry (2004), 125(4), 553-560

CODEN: JFLCAR; ISSN: 0022-1139

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 141:349790

A series of reactions of SF5CF2CF2I and SF5(CF2)4I with F2C:CF2 was carried out in an effort to find the most effective methods for

chain-extension. SF5(CF2)8I and SF5(CF2)10I were prepared and isolated. The reaction conditions for the addition of H2C:CH2 were also investigated.

The crystal structure of SF5(CF2)4CH2CH2I was determined

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 5 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:292782 CAPLUS

DOCUMENT NUMBER: 135:121980

TITLE: A simple procedure for nucleophilic

perfluoroalkylation of organic and inorganic

substrates

AUTHOR(S): Petrov, V. A.

CORPORATE SOURCE: Experimental Station, Central Research and

Development, E. I. Du Pont de Nemours and Co.,

Wilmington, DE, 19880-0328, USA

SOURCE: Tetrahedron Letters (2001), 42(19), 3267-3269

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 135:121980 The mixture perfluoroalkyl iodide and

> tetrakis (dimethylamino) ethylene is used for the nucleophilic perfluoroalkylation. The reaction of chlorotrimethylsilane with

perfluoroalkyl iodide and tetrakis (dimethylamino)

ethylene in diglyme gives perfluoroalkyltrimethylsilane in 55-81%

yield. The interaction of this system with organic electrophiles such as benzoyl and benzensulfonyl chlorides, aliphatic and aromatic aldehydes and activated ketones leads to the formation of the corresponding condensation products in 24-62% yield.

REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 6 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:180342 CAPLUS

DOCUMENT NUMBER: 134:366988

TITLE: Reactions of Halofluorocarbons with Group 6 Complexes

> M(C5H5)2L (M = Mo, W; L = C2H4, CO). Fluoroalkylation at Molybdenum and Tungsten, and at Cyclopentadienyl or

Ethylene Ligands

AUTHOR(S): Hughes, Russell P.; Maddock, Susan M.; Guzei, Ilia A.;

Liable-Sands, Louise M.; Rheingold, Arnold L.

Departments of Chemistry Burke Chemistry Laboratory, CORPORATE SOURCE:

Dartmouth College, Hanover, NH, 03755-3564, USA

SOURCE: Journal of the American Chemical Society (2001),

123(14), 3279-3288

CODEN: JACSAT; ISSN: 0002-7863

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 134:366988

The Mo(II) and W(II) complexes [MCp2L] (Cp = η 5-cyclopentadienyl; L = C2H4, CO) react with perfluoroalkyl iodides to give a variety of products. The Mo(II) complex [MoCp2(C2H4)] reacts with perfluoro-BuI or perfluorobenzyl iodide with loss of ethylene to give the 1st examples of fluoroalkyl complexes of Mo(IV), MoCp2(CF2CF2CF3)I (8) and MoCp2(CF2C6F5)I (9), one of which (8) was crystallog. characterized. In contrast, the CO analog [MoCp2(CO)] reacts with perfluorobenzyl iodide without loss of CO to give the crystallog. characterized salt, [MoCp2(CF2C6F5)(CO)]+I- (10), and the W(II) ethylene precursor [WCp2(C2H4)] reacts with perfluorobenzyl iodide without loss of ethylene to afford [WCp2(CF2C6F5)(C2H4)]+I-(11). These observations demonstrate that the metal-C bond is formed 1st. In further contrast the W precursor [WCp2(C2H4)] reacts with perfluoro-BuI, perfluoro-iso-Pr iodide, and pentafluorophenyl iodide to give fluoroalkyl- and fluorophenyl-substituted cyclopentadienyl complexes WCp(η 5-C5H4RF)(H)I (12, RF = CF2CF2CF2CF3; 15, RF = CF(CF3)2; 16, RF = C6F5); the Mo analog MoCp(η 5-C5H4RF)(H)I (14, RF = CF(CF3)2) was obtained in similar fashion. The W(IV) hydrido compds. react with iodoform to afford the corresponding diiodides $WCp(\eta 5-C5H4RF)12$ (13, RF = CF2CF2CF3; 18, RF = CF(CF3)2; 19, RF = C6F5), two of which (13 and 19) were crystallog. characterized. The carbonyl precursors [MCp2(CO)] each react with perfluoro-iso-Pr iodide without loss of CO, to afford the exo-fluoroalkylated cyclopentadiene M(II) complexes MCp(n4-C5H5RF) (CO)I (21, M = Mo; 22, M = W); the exo-stereochem. for the fluoroalkyl group is confirmed by an x-ray structural study of 22. ethylene analogs [MCp2(C2H4)] react with perfluoro-tert-BuI to yield the products MCp2[CH2CH2C(CF3)3]I (25, M = Mo; 26, M = W) resulting from fluoroalkylation at the ethylene ligand. Attempts to provide pos. evidence for fluoroalkyl radicals as intermediates in reactions of primary and benzylic substrates were unsuccessful, but trapping expts. with CH3OD (to give RFD, not RFH) indicate that fluoroalkyl anions are the intermediates responsible for ring and ethylene fluoroalkylation in the reactions of secondary and tertiary fluoroalkyl substrates.

ANSWER 7 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

65

ACCESSION NUMBER: 2000:200696 CAPLUS

DOCUMENT NUMBER: 133:17879

TITLE: Controlled step-wise telomerization of vinylidene

fluoride, hexafluoropropene and trifluoroethylene with

THERE ARE 65 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

iodofluorinated transfer agents

AUTHOR(S):

REFERENCE COUNT:

Balague, J.; Ameduri, B.; Boutevin, B.; Caporiccio, G. CORPORATE SOURCE: Ecole Nationale Superieure de Chimie, UMR 5076 (CNRS),

Laboratory of Macromolecular Chemistry, Montpellier,

34296, Fr.

SOURCE: Journal of Fluorine Chemistry (2000), 102(1-2),

253-268

CODEN: JFLCAR; ISSN: 0022-1139

PUBLISHER: Elsevier Science S.A. DOCUMENT TYPE: Journal LANGUAGE: English

Highly fluorinated cotelomers having the structure F(TFE)w(VDF)x(HFP)y(TrFE)zI containing one or several tetrafluoroethylene (TFE), vinylidene fluoride (or 1,1-difluoroethylene, VDF), hexafluoropropene (HFP) or trifluoroethylene (TrFE) base units were synthesized by thermal step-wise cotelomerization of these fluoroolefins with perfluoroalkyl iodides. 1H and 19F NMR allowed one to characterize these cotelomers and to assess the defects of chaining and the mol. wts. While the monoadduct produced from VDF exclusively exhibits RFCH2CF2I structure, that prepared from TrFE was composed of RFCFHCF2I and RFCF2CFHI isomers, the ratio of which is directed from the electrophilicity of RF√ radical. The reactivity of the C-I bond in RF-Q-CXY-I depends on the nature of the Q spacer and on the reactivity of the fluorinated monomer (e.g., thermal initiations of VDF, TrFE and HFP were efficient from 180° , 195° and 210° , resp.). The mechanism of the addition of the radical generated from the iodinated transfer agent to the fluoroalkene is explained by means of its electrophilic attack to the more nucleophilic (i.e., the less electrophilic) side of the olefin. Ethylenation of these fluorocotelomers was successfully achieved from various initiations (thermal, redox or from peroxides) with best results from redox catalysis. Thermal properties of several fluorotelomers (glass transition temps., Tg and melting temps., Tm) were assessed. They were linked to the number of consecutive CF2 groups (for the crystalline zones) and bulky side groups which induced amorphous regions.

REFERENCE COUNT: 80 THERE ARE 80 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 8 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:200694 CAPLUS

DOCUMENT NUMBER: 133:17205

TITLE: Diiododifluoromethane: an excellent telogen for the

preparation of 1,3-diiodofluoropropane derivatives

AUTHOR(S): Yang, Z.-Y.

CORPORATE SOURCE: Experimental Station, Central Research & Development,

E. I. Du Pont de Nemours and Co., Wilmington, DE, USA

SOURCE: Journal of Fluorine Chemistry (2000), 102(1-2),

239-241

CODEN: JFLCAR; ISSN: 0022-1139

PUBLISHER: Elsevier Science S.A.

DOCUMENT TYPE: Journal LANGUAGE: English

AB CF2I2 reacted with olefins thermally to give 1,3-diiodofluoropropane derivs. Both fluorinated and non-fluorinated alkenes gave good yields of adducts. Reaction with ethylene, propylene,

perfluoroalkylethylene, vinylidene fluoride, and trifluoroethylene provided the corresponding adducts in 58-86% yields. With tetrafluoroethylene, 1-to-1 adduct was predominantly formed along with small amts. of higher homologs. In contrast to **perfluoroalkyl** iodides, CF2I2 also readily added to perfluorovinyl ethers to give

1,3-diiodoperfluoroethers.

REFERENCE COUNT: 29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 9 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:403224 CAPLUS

DOCUMENT NUMBER: 127:81545

TITLE: Selective Fluoroalkylation of Cyclopentadienyl and

Ethylene Ligands in Reactions of

Perfluoroalkyl Iodides with Low-Valent Complexes of Molybdenum and Tungsten: Evidence for a Fluorocarbanion Mechanism

Hughes, Russell P.; Maddock, Susan M.; Rheingold,

Arnold L.; Liable-Sands, Louise M.

CORPORATE SOURCE: Department of Chemistry 6128 Burke Laboratory,

Dartmouth College, Hanover, NH, 03755, USA

SOURCE: Journal of the American Chemical Society (1997),

119(25), 5988-5989

CODEN: JACSAT; ISSN: 0002-7863

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 127:81545

AB Reactions of perfluoroalkyl iodide ICF(CF3)2 with

[MCp2(C2H4)] (2, M = Mo; 3, M = W) afford the hydrido complexes
[{C5H4CF(CF3)2}(Cp)]MHI (4, M = Mo; 5, M = W) quant. The structure of 5
has been confirmed by x-ray crystallog.; monoclinic, P 31/c, a = 6.268(2),

b = 33.748(6), c = 7.616(2) Å, $\beta = 112.01(2)$ °, V =

1493.6(6) Å3, Z = 4. In contrast, the tertiary perfluoroalkyl iodide IC(CF3)3 reacts with 2 or 3 cleanly at the ethylene

ligand to give 6 and 7. X-ray crystallog. also confirms the structure of 7; monoclinic, P21/n, a = 12.8936(1), b = 7.557(1), c = 38.9142(2) Å, β = 95.0237(2)°, V = 3777.46(6) Å3, Z = 8 (two chemical

identical, crystallog. independent mols.). Running these reactions in the presence of radical traps like perdeuterotoluene or dihydroanthraene provides no evidence for fluoroalkyl radical intermediates, but pos. signatures for fluorocarbanion intermediates are provided by trapping expts. with CH3OD, and by observation of fluoroolefins F2C:CF(CF3) and F2C:C(CF3)2 among the reaction products from ICF(CF3)2 or IC(CF3)3 resp.

REFERENCE COUNT: 27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 10 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:650376 CAPLUS

DOCUMENT NUMBER:

123:256153

TITLE:

AUTHOR(S):

Preparation of perfluoroalkyl

iodides from perfluorocarboxylic acids

INVENTOR(S):
Ueda, Kunimasa

PATENT ASSIGNEE(S): Idemitsu Petrochemical Co, Japan SOURCE: Jpn. Kokai Tokkyo Koho, 12 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 07109234	A2	19950425	JP 1993-277348	19931012
PRIORITY APPLN. INFO.:			JP 1993-277348	19931012

OTHER SOURCE(S): MARPAT 123:256153

AB In preparation of CnFmI (n ≥ 1; 3 ≤ m ≤ 2n + 1), useful as perfluoroalkylating agents by treatment of CnFmCO2H with I and Bz2O2 (I) in an organic solvent, a homogeneous solution or suspension of I in the organic solvent, obtained by treatment of I containing H2O with the organic solvent, which is suitable for the above iodination and practically immiscible with H2O, to extract I followed by separation and removal of the aqueous layer, is

I and the solvent. The method is prevented from danger of explosive I. I containing 25 weight% H2O was suspended in C1(CH2)4Cl (II) and the upper H2O layer was removed. The suspension was treated with CF3(CF2)7CO2H, II, and I under reflux at 100° for 1 h to give 77% CF3(CF2)7I.

L7 ANSWER 11 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:321022 CAPLUS

DOCUMENT NUMBER:

123:313360

TITLE:

Free-radical addition of 2-

(perfluoroalkyl)ethanethiols to alkenes, alkadienes, cycloalkenes, alkynes and vinyl monomers. [Erratum to

document cited in CA119:225554]

AUTHOR(S): B:

Brace, Neal O.

CORPORATE SOURCE: Wheaton College, Wheaton, IL, 60187, USA

SOURCE: Journal of Fluorine Chemistry (1995), 70(1), 145

CODEN: JFLCAR; ISSN: 0022-1139

PUBLISHER: Elsevier DOCUMENT TYPE: Journal LANGUAGE: English

AB The errors were not reflected in the abstract or the index entries.

L7 ANSWER 12 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:136059 CAPLUS

DOCUMENT NUMBER: 120:136059

TITLE: Perfluoroalkyl halides and derivatives as precursors

for oil and water repellants and surfactants

INVENTOR(S): Behr, Frederick E.; Dams, Rudolf J.; DeWitte, Johan

E.; Hagen, Donald F.

PATENT ASSIGNEE(S): Minnesota Mining and Manufacturing Co., USA

SOURCE:

Can. Pat. Appl., 67 pp.

CODEN: CPXXEB

DOCUMENT TYPE: Patent

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CA 2071596	AA	19930111	CA 1992-2071596	19920618
EP 526976	A1	19930210	EP 1992-305710	19920622
EP 526976	B1	19970115		
R: BE, CH, DE,	FR, GB	, IT, LI, NL		
JP 05345732	A2	19931227	JP 1992-183345	19920710
JP 3231844	B2	20011126		
JP 2002138078	A2	20020514	JP 2001-204928	19920710
PRIORITY APPLN. INFO.:			US 1991-728184 A	19910710
			JP 1992-183345 A	3 19920710

OTHER SOURCE(S): MARPAT 120:136059

The title compds. comprise a mixture of straight and branched perfluoroalkyl groups bonded to Cl, Br, or I through a F-free alkylene group. Perfluorodecyltetrahydroiodide (prepared from perfluorosulfonyl fluoride, 40% straight and 60% branched, treated first with I, then with C2H4) was derivatized to thiol functionality by treatment with thiourea in EtOH to give perfluorodecyltetrahydrothiol (I). I was added to a reaction mixture containing hexamethoxymethylmelamine to give a I-melamine condensate (II, 1:4 mol ratio). A 50/50 polyester/cotton fabric blend was treated with an emulsion of II at 0.3%, dried and cured at 150°, to give a fabric with oil resistance (AATCC 118-1975) 5 and 5 after 1 dry cleaning, vs. 3 and 2, resp., for a precursor perfluorodecyltetrahydroiodide having all straight chain perfluoroalkyl groups.

L7 ANSWER 13 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1993:625554 CAPLUS

DOCUMENT NUMBER: 119:225554

TITLE: Free-radical addition of 2-

(perfluoroalkyl) ethanethiols to alkenes, alkadienes,

cycloalkenes, alkynes and vinyl monomers

AUTHOR(S): Brace, Neal O.

CORPORATE SOURCE: Wheaton Coll., Wheaton, IL, 60187, USA

SOURCE: Journal of Fluorine Chemistry (1993), 62(2-3), 217-41

CODEN: JFLCAR; ISSN: 0022-1139

DOCUMENT TYPE: Journal LANGUAGE: English

AB The free-radical addition of 2-(perfluoroalkyl)ethanethiols (RFCH2CH2SH) to alkenes, cycloalkenes, alkadienes and alkynes has been studied to determine (1) the mode of reaction, i.e., the stereochem., regiochem. and any skeletal changes; (2) the relative reactivity towards unsats. of various structures and classes as affected by the presence of the RF group; and (3) the influence of the reaction conditions on the rate of addition or selectivity for different products. Adducts from 2-(F-hexyl)ethanethiol (1) and alkenes have been obtained in high yield, but containing small amts. of regio isomers. For example, compound 1 with 1-heptene gave 96%
1-[2-(F-hexyl)ethanethio]heptane, as well as 0.61% 2- and 2.22%
3-[2-(F-hexyl)ethanethio]heptane. Dienes gave chiefly linear adducts adducts; small amts. of cyclic isomers were also formed. Compound 1 added readily with free-radical initiation to vinyl monomers such as styrene and vinyl acetate, and to phenylacetylene, propargyl acetate and Et propynoate.

L7 ANSWER 14 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1988:454260 CAPLUS

DOCUMENT NUMBER: 109:54260

TITLE: Synthesis and chemistry of perfluoro-2-iodo-2-

methylalkanes

AUTHOR(S): Probst, A.; Raab, K.; Ulm, K.; Von Werner, K.

CORPORATE SOURCE: Werk Gendorf, Hoechst A.-G., Burgkirchen, D-8269, Fed.

Rep. Ger.

SOURCE: Journal of Fluorine Chemistry (1987), 37(2), 223-45

CODEN: JFLCAR; ISSN: 0022-1139

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 109:54260

AB CF3(CF2)nC(CF3)2I (I, n = 1, 2) were obtained from CF3(CF2)nCF:C(CF3)2 (n = 0, 1) by formal addns. of IF that required substantial alterations of known procedures. I (n = 1, 2) are the most reactive alkyl halides known so far, and they are also very toxic. The reactions studied included (a) Nucleophilic attack of anions at the iodine, leading to perfluoroalkenes, (b) elimination of IF, caused by metals or metal complexes, (c) pyrolysis, to selectively give perfluoroisobutene and n-perfluoroalkyl iodides, (d) photolysis, and (e) thermally induced insertions into the carbon-iodine bond. Screening results on the inhalation toxicity of the iodides and of some other fluoro-compds. are also reported.

L7 ANSWER 15 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1987:33621 CAPLUS

DOCUMENT NUMBER: 106:33621

TITLE: ω -Perfluoroalkyl- α -olefin polymers

INVENTOR(S): Su, Aaron Chung Liong

PATENT ASSIGNEE(S): du Pont de Nemours, E. I., and Co., USA

SOURCE: Eur. Pat. Appl., 18 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 193202	A2	19860903	EP 1986-102650	19860228

EP 193202 A3 19870722 R: DE, FR, GB, IT US 4617363 19861014 US 1985-707013 Α 19850228 CA 1275544 A1 19901023 CA 1986-502693 19860225 JP 61204208 A2 19860910 JP 1986-42027 19860228 JP 03080804 B4 19911226 JP 01103609 A2 19890420 JP 1988-208460 19880824 PRIORITY APPLN. INFO.: US 1985-707013 A 19850228 Polymers of CH2:CH(CH2)nR (R = C2-10 perfluoroalkyl; n = 2-8) have high gas-selective permeability and solvent resistance. Thus, 20 mL C4F9(CH2)2CH:CH2 (prepared by addition of C4F9(CH2)2I and C2H4, then dehydroiodination with tert-BuOK), 3 mmol iso-Bu3Al, and 0.3 g TiCl4 on MgCl2 were heated at 50° for 4 h to form poly(4-perfluorobuty1-1butene) having 100% modulus 3.4 MPa and tensile strength 10.3 MPa, which was insol. in hydrocarbon, fluorocarbon, and polar solvents, and showed CO2 permeability (80:20 CO2/CH4) 7.7 + 10-8 cm3-cm/cm2-s-cm Hg and selectivity [(CO2/CH4 in product)/(CO2/CH4 in feed)] 9; vs. 1.2 + 10-9 cm3-cm/cm2-s-cm Hg and 10 for Teflon; or 5 + 10-8cm3-cm/cm2-s-cm Hg and 5 for natural rubber.

L7 ANSWER 16 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1987:17597 CAPLUS

DOCUMENT NUMBER: 106:17597

TITLE: Fluoroiodo compounds

PATENT ASSIGNEE(S): du Pont de Nemours, E. I., and Co., USA

SOURCE: Jpn. Kokai Tokkyo Koho, 11 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 61134329	A2	19860621	JP 1985-269194	19851129
US 4650913	Α	19870317	US 1984-676100	19841129
EP 194348	A2	19860917	EP 1985-115132	19851129
EP 194348	A3	19870819		
EP 194348	B1	19900124		
R: CH, DE, FR,	LI			
PRIORITY APPLN. INFO.:			US 1984-676100 A	19841129

OTHER SOURCE(S): CASREACT 106:17597; MARPAT 106:17597

AB RCR1R2CR3R4I (R = C1-18 perfluorohydrocarbon residue; R1-4 = H, C1-16 hydrocarbon, mono- or bicyclic aliphatic residue) are prepared by addition of R1R2C:CRR4 with RI in the presence of R5SO2M (R5 = C1-18 hydrocarbon residue, M = alkali metal, NH4, etc.) as initiators. Thus a mixture of norbornene 0.1, CF3(CF2)7I 0.02, and PhSO2Na 0.03 mol in DMF was stirred under Ar to give 93% 2-iodo-3-(perfluorooctyl)norbornane.

L7 ANSWER 17 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1985:614859 CAPLUS

DOCUMENT NUMBER: 103:214859

TITLE: Fluoroalkyl-substituted iodoalkanes

INVENTOR(S):
Von Werner, Konrad

PATENT ASSIGNEE(S): Hoechst A.-G., Fed. Rep. Ger.

SOURCE: Ger. Offen., 19 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 3338300	A1	19850502	DE 1983-3338300	19831021
EP 140254	A1	19850508	EP 1984-112300	19841012
EP 140254	B1	19881005		
R: BE, CH, DE,	FR, GB	, IT, LI, NL		
JP 60106533	A2	19850612	JP 1984-218786	19841019
us 4587366	Α	19860506	US 1984-663084	19841019
PRIORITY APPLN. INFO.:			DE 1983-3338300 A	19831021
OTHER SOURCE(S):	MARPAT	103:214859		

(Fluoroalkyl)alkyl iodides were prepared by reaction of a fluorinated alkyl iodide with warm (un) substituted alkenes in the presence of a finely-divided metal catalyst chosen from elements with atomic nos. 24-30, 42-48, or 74-79. I(CF2)4I (0.68 mol) reacted with G0.68 mol CH2:CH2 in the presence of Ru at 170°/2 MPa in 33 h to give 97.8% yield of I(CH2)2(CF2)4(CH2)2I with 98.5% conversion I(CF2)4I.

ANSWER 18 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1984:51074 CAPLUS

DOCUMENT NUMBER:

100:51074

TITLE:

Branched 1,2-bis(perfluoroalkyl)ethenes

INVENTOR(S):

Riess, Jean Georges; Jeanneaux, Francois; Le Blanc,

Maurice; Lantz, Andre

PATENT ASSIGNEE(S):

Produits Chimiques Ugine Kuhlmann, Fr.

SOURCE:

Eur. Pat. Appl., 12 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent French

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

P.A	TENT NO.		1	KIND	DATE	APPLICATION NO.		DATE
			-					
E	90712			A 1	19831005	EP 1983-400569		19830318
EF	90712			В1	19850619			
	R: AT,	BE,	CH, I	DE, F	R, GB, IT,	LI, NL, SE		
FF	2523956			A1	19830930	FR 1982-5165		19820326
FF	2523956			В1	19850524			
US	4613708			Α	19860923	US 1983-467648		19830218
ΓA	13872			E	19850715	AT 1983-400569		19830318
AU	8312872			A1	19830929	AU 1983-12872		19830325
AU	554646			B2	19860828			
JE	58174334			A2	19831013	JP 1983-49089		19830325
JE	01055251			B4	19891122			
ES	521022			A1	19840601	ES 1983-521022		19830325
C.F	1186344			A1	19850430	CA 1983-424516		19830325
PRIORIT	Y APPLN.	INFO.	:			FR 1982-5165	Α	19820326
						EP 1983-400569	Α	19830318

OTHER SOURCE(S): CASREACT 100:51074; MARPAT 100:51074

Perfluoroalkyl iodides reacted with

mono(perfluoroalkyl)ethylenes, and the addition products obtained were dehydriodinated to yield RCH: CHR1 (R and R1 are perfluoroalkyl and one or both of them is/are branched), useful in emulsions. A mixture of CH2:CHCF(CF3)2 and (CF3)2CFI was heated at 200° to give (CF3)2CFCH2CHICF(CF3)2, which was treated with KOH in EtOH to give (CF3) 2CFCH: CHCF(CF3) 2.

ANSWER 19 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1982:217164 CAPLUS

DOCUMENT NUMBER:

96:217164

TITLE:

Synthesis of fluorinated acetylenes

AUTHOR(S):

Baum, Kurt; Bedford, Clifford D.; Hunad, Ronald J.

CORPORATE SOURCE: Fluorochem, Inc., Azusa, CA, 91702, USA

SOURCE: Journal of Organic Chemistry (1982), 47(12), 2251-7

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 96:217164

AB New routes to fluorinated acetylenes were developed based on addns. of iodofluorocarbons to silylacetylenes. Free radical addition of α ω-diiodoperfluoroalkanes to (trimethylsilyl)acetylene gave the iodotrimethylsilyl olefins Me3SiIC:CH(CF2)nCH:CISiMe3, which reacted with KOCMe3 or 1,8-diazabicyclo[5,4,0]undec-7-ene to give (trimethylsilyl)acetylenes and, with an excess of the base, the free diacetylenes. Perfluoroalkyl iodides similarly gave (perfluoroalkyl)acetylenes. The addition of perfluoroheptyl iodide to PhC.tplbond.CH followed by treatment with KOCMe3 gave 1-phenylperfluorononyne. The peroxide-catalyzed reaction of perfluoroalkyl iodides (RFI) and Me3SiC.tplbond.CSiMe3 gave 1:1 adducts, RF(Me2SiCH2)IC:CHSiMe3, resulting from intramol. H abstraction by the initially formed vinyl radical. The thermal reaction

of **perfluoroalkyl iodides** and diiodides with bis(trimethylsilyl)acetylene in the presence of free iodine gave the (trimethylsilylacetylenes, which were desilylated with KF. A route to diacetylenes was investigated based on addition of perfluoroiodo compds. to

ethylene, dehydroiodination, brominations, and eliminations.

L7 ANSWER 20 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1978:512054 CAPLUS

DOCUMENT NUMBER: 89:112054

TITLE: Modification of fiber surfaces by monomeric additives.

Part II. Absorption of fluorocarbon additives by

poly(ethylene terephthalate)
Mares, F.; Oxenrider, B. C.

CORPORATE SOURCE: Chem. Res. Cent., Allied Chem. Corp., Morristown, NJ,

USA

SOURCE: Textile Research Journal (1978), 48(4), 218-29

CODEN: TRJOA9; ISSN: 0040-5175

DOCUMENT TYPE: Journal LANGUAGE: English

AUTHOR(S):

AB Evaluation of 3 different methods for the modification of poly(
ethylene terephthalate) fiber surfaces showed that coating the

fibers with reactive group-containing additives from a dioxane solution followed

by annealing produces the best results. The additives, e.g., an isomeric mixture of dihydrogen bis (4-perfluoroisopropoxy-3,3,4,4-tetrafluorobutyl pyromellitates and mono (4-perfluoroisopropoxy-3,3,4,4-tetrafluorobutyl) phthalate [62478-07-3], were not simply deposited on the fiber surfaces, as demonstrated by scanning electron microscopy, but penetrated the fiber and interacted with the polymer matrix to form oligomers near the fiber surface. Additive concns. as low as 0.25% produced excellent water and oil repellency that was durable to washing and dry cleaning owing to the insoly. of the additives in soap solns. and dry cleaning solvents. The modified fibers could be knitted and dyed without affecting the surface properties.

L7 ANSWER 21 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1978:74021 CAPLUS

DOCUMENT NUMBER: 88:74021

TITLE: Addition to gaseous nonhalogenated olefins and

acetylenes of perfluoroalkyl iodides

INVENTOR(S):
Knell, Martin

PATENT ASSIGNEE(S): Ciba-Geigy Corp., USA

SOURCE: U.S., 7 pp.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4058573	Α	19771115	US 1974-457879	19740404
PRIORITY APPLN. INFO.:			US 1967-693148	A2 19671226
			US 1970-4179	A2 19700114
			US 1971-159515	A1 19710702

AB A gaseous olefin (e.g., ethylene, propene, and 2-butene) or acetylene was bubbled through liquid perfluoroalkyl iodide [e.g., 1-iodoperfluoroheptane (I) and 1-iodoperfluorohexane] or a mixture of perfluoroalkyl iodides at .apprx.50-100° in the presence of a free radical generating catalyst to give the corresponding adducts, essentially free of telomers. Thus, ethylene was bubbled through a mixture of I and Bz2O2 at 80-5° for .apprx.10.5 h to give 1,1,2,2-tetrahydro-1-iodoperfluorononane (88.5% yield) free of telomers. About 18 other fluoroiodoalkanes or -alkenes were similarly prepared

ANSWER 22 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1974:48569 CAPLUS

DOCUMENT NUMBER: 80:48569

TITLE: Fluoroalkyl iodide telomers

Rudolph, Werner; Massonne, Joachim INVENTOR(S):

Kali-Chemie A.-G. PATENT ASSIGNEE(S): SOURCE: Ger. Offen., 11 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2218451	A 1	19731031	DE 1972-2218451	19720417
CH 580552	Α	19761015	CH 1973-4264	19730323
GB 1415245	Α	19751126	GB 1973-16199	19730404
NL 7304737	Α	19731019	NL 1973-4737	19730405
BE 798268	A1	19731016	BE 1973-130056	19730416
FR 2180863	A 1	19731130	FR 1973-13754	19730416
JP 49014407	A2	19740207	JP 1973-43040	19730416
IT 982785	Α	19741021	IT 1973-23107	19730417
PRIORITY APPLN. INFO.:			DE 1972-2218451 A	19720417

AB Fluoroalkyl iodide telomers were prepared by 1:1 addition of ethylene to C6-12 perfluoroalkyl iodide telomers (I) in the presence of peracid catalysts, e.g. peracetic acid [79-21-0] at 70-100.deg.. Thus, 40 standard 1. ethylene/hr and 30 g catalyst solution/hr (11.6% HO2Ac in C2F3Cl3) were passed into a reactor containing 4000 g

I 8 hr at 70-80.deg. to give 98.3% monoethylene adduct of I.

ANSWER 23 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1974:14545 CAPLUS

DOCUMENT NUMBER: 80:14545

TITLE: Perfluoroalkyl iodides INVENTOR(S): Oda, Yoshio; Suhara, Manabu

PATENT ASSIGNEE(S): Asahi Glass Co., Ltd.

SOURCE: Jpn. Kokai Tokkyo Koho, 4 pp. CODEN: JKXXAF

DOCUMENT TYPE:

Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	-			
JP 48052706	A2	19730724	JP 1971-88200	19711108
JP 56002054	В4	19810117		

PRIORITY APPLN. INFO.: JP 1971-88200 A 19711108

Catalytic action of Mo or W fluoride gave perfluoroalkyl iodides from C2F4, iodine, and IF5. E.g., 20.3 g C2F4 was fed to a mixture of iodine 16.8 g, IF5 7.4 g, and MoF6 100 mg at 75° in 2.8 hr at 4-12 kg/cm2 to give 39.1 g pentafluoroethyl iodide.

ANSWER 24 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1969:501287 CAPLUS

DOCUMENT NUMBER: 71:101287

TITLE: Iodoperfluoroalkanes

INVENTOR(S): Knell, Martin

PATENT ASSIGNEE(S): Geigy, J. R., A.-G. Ger. Offen., 11 pp. SOURCE:

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
		C3		DE 1968-1816706	19681223
	CH 509954	Α	19710715	CH 1968-509954	19681202
	GB 1210730	Α	19701028	GB 1968-1210730	19681224
	FR 1597171	Α	19700622	FR 1968-1597171	19681226
PRIC	RITY APPLN. INFO.:			US 1967-693148 A	19671226
AB	Perflouroalkyl iodi	ides [C	xF2x+1] (x =	4-14), which are liquid	at reaction
	temperature, are ac	ded to	gaseous olef	ins and acetylenes in t	he presence of a
	catalyst giving fre	e radio	als, at atmo	spheric or lower pressu	re and
	50-220°. Thus, eth				
				by weight 1-iodoperfluo	roheptane
				to give 88.5% 1,1,2,2-t	
				5°. Similarly prepared	
				Eluorodecane, b. 119-20°	
		_	_	prononane, b. 82-4°;	,
				prononane, b. 86-90°;	
	1,1,2,2-tetrahydro-				
	1,2-dihydro-1-iodor			me, b. 107-10 ,	
	1,2 dinyd10-1-10d0p	erridor	ononene.		

ANSWER 25 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1969:402955 CAPLUS

DOCUMENT NUMBER: 71:2955

TITLE: Perfluoroalkyl fluoriodides and their use for

telomerization of perfluoroalkyl

iodides with olefins

Rondestvedt, Christian S., Jr. INVENTOR(S): PATENT ASSIGNEE(S): du Pont de Nemours, E. I., and Co.

SOURCE: Fr., 14 pp. CODEN: FRXXAK

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT:

PATENT NO. KIND DATE APPLICATION NO. -----19680419 FR 1967-104939 FR 1521775 19670502 PRIORITY APPLN. INFO.: US The title preparation and telomerization, especially telomerization of tetrafluoroethylene (I) is described. Thus, 4.67 g. liquid C1F3 was added with stirring at -70° to a solution of 33.4 g. n-C6F13I in 100 ml. perfluorohexane. The temperature raised by exothermic reaction to 0°, C6F14 and excess C6F13I were evaporated in vacuo <0° to give C6F13IF2. Similarly, C2F5I and C1F3 gave C2F5IF2 with some C2F5IF4; 6 moles n-C4F9I with 1 mole C1F3 gave n-C4F9IF2, m. 150°; 0.0337 mole n-C10F21I with 0.03 mole ClF3 gave Cl0F21IF2, solid. Addition of 40.7 g. liquid ClF3 to 206 g. n-C4F9I in 2614 g. C6F14 at -80° gave a precipitate, dissolved by slow heating to -30° ; the solution was cooled to -55° , and more 69.3 q. ClF3 was added to give 80% n-C4F9IF4 m. 10° (stable at -20° , but decomposed at room temperature). Dry He, saturated at 0° with C1F3 vapor, was bubbled in C2F5I at 70° to absorb 2 moles C1F3, and the mixture was evaporated in vacuo <0° to give C2F5IF2, m. 18° By the He-method, the reaction of 0.1 mole ClF3 with 0.5 mole (CF3)2CFI at -70° gave 99% (CF3)2CFIF2, pasty solid; 0.1 mole ClF3 with 0.5 mole C6F13I gave n-C6F13IF2, m. 35°; 25 g. CF3I in 30 ml. C6F14 with He-ClF3 at -80 to 30° gave a solid, decomposing 0°. A mixture of 42.6 g. C2F5I and 7 g. BrF5 did not react at -50°; on heating, a violent reaction gave C2F5IF2 and C2F5IF4. A similar reaction of 0.15 mole n-C4F9I with 0.06 mole BrF5 at -80 to -30° gave an equimolar mixture of n-C4F9-IF2 and C4F9IF4, m. 18°. The mixture of 0.1 mole n-C4F9I in 146 g. C6F14 with 0.04 mole BrF5 at -80 to 0° gave a mixture of 64% n-C4F9IF4 and 36% n-C4F9IF2, m. 16°. A similar reaction of C4F9I with BrF3 at the same conditions gave the same compds. For the telomerization, 5.9 g. C6F13IF2 and 100 g. n-C3F7I in a glass autoclave in vacuo was heated to 70°, and 33.4 g. I was introduced at $70-8^{\circ}/8.4$ bar, in 10 min. After 32 min. at 70° , the pressure fell to 2.25 bar, to give a mixture of telomers F(CF2)nI with predominately odd n from 3 to 19. Other methods for telomerization of I are related, giving telomers with n 0-7, 1-6, 3-11, 2-20, with predominately odd or even n, using as telogen C4F9I and C2F5I, and as catalyst C4F9IF2 and C3F7IF4. RIF2 are more active as catalysts than RIF4. The telomerization of ethylene with n-C4F9I, catalyzed with n-C4F9IF2, at 60-80° gave a low yield of n-C4F9CH2CH2I and traces of telomers with n = 2,3,4. Similarly, vinyl fluoride gave n-C4F9CH2CHFI, and vinylidene fluoride gave n-C4F9CH2CF2I, b760 120°.

L7 ANSWER 26 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1966:43304 CAPLUS

DOCUMENT NUMBER: 64:43304

ORIGINAL REFERENCE NO.: 64:8031h,8032a-c

TITLE: Perfluoroalkyl iodides

INVENTOR(S): Blanchard, Wesley A.; Rhode, Judson C.

PATENT ASSIGNEE(S): E. I. du Pont de Nemours & Co.

SOURCE: 4 pp.
DOCUMENT TYPE: Patent
LANGUAGE: Unavailable

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 3226449 19651228 US 1962-200299 19620606

AB Perfluoroalkyl iodides containing 6-12 C atoms may be

prepared by injecting C2F4 and a free radical-generating catalyst into a mixture of (1) a **perfluoroalkyl iodide** chosen from C2F5I and mixts. of C2F5I and C4F9I and (2) a portion of free radical generating catalyst. The mixture is heated to 80-170° under a pressure of 225-700 psi. The injection of the C2F4 mixture is continued until there are 0.25-1.2 moles of C2F4 per mole of **perfluoroalkyl iodide**. The amount of catalyst should be 0.25-0.9 weight-% of the total reaction mixture Product yields of 70% or better are obtained. The preferred catalyst is di-tert-butyl peroxide. The preferred reaction temperature with

the

catalyst is 165°. The preferred mole ratio of C2F4-iodide is 0.4-1.1. The table gives yields obtained under various conditions. The products may be used as intermediates for preparing oil and water repellants for textiles and as intermediates for the preparation of perfluoroalkanecarboxylic acids which are surface active agents. Parts, 2F6I, C4F9I, Total tert-Bu2O2used, weight %, Temperature, Ratio of C2F4 added

to

initial iodide moles, % yield C6F13I-C12F25I, Based on C2F4, Based on iodide; 800, 800, 0.9, 165°, 0.654, 93.9, 74.5; 1200, 400, 0.89, 165°, 0.961, 98, 80.1; 1200, 400, 0.78, 165°, 0.67, 98+, 73.6; 1200, 400, 0.85, 165°, 0.767, 90.7, 81.9; 1200, 400, 0.91, 165°, 0.86, --, 60; 1200, 400, 0.78, 175°, 1.2, 65, 65;

L7 ANSWER 27 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1966:18698 CAPLUS

DOCUMENT NUMBER: 64:18698

ORIGINAL REFERENCE NO.: 64:3349g-h,3350a-b
TITLE: Perfluoroalkyl iodides

INVENTOR(S):

Bloechl, Walter

SOURCE:
DOCUMENT TYPE:

13 pp. Patent

LANGUAGE:

Unavailable

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	NL 6414504		19650614	NL 1964-14504	19641211
PRIC	DRITY APPLN. INFO.:			DE	19631212
AB				yields (without formi	
				with a saturated haloal?	
				0-350° for <10 min. The	
				<pre>fluoroheptyl iodide (I)</pre>	
	CF2:CF2 (II) was fe	ed conti	inuously (0.1	g. I/min.) into a Pyro	ex glass tube
				ime of 3 min., the mix	
				uid formed was distille	
				'l iodide (III), b760 1	78°,
				due, containing 90%	
	perfluoroundecyl ic				
				Similarly, CF3CF2I (V	
				330°, and the product	
				1-iodononafluorobutane	
				oheptadecafluorooctane	
				erfluoro compds. prepar	
				reaction conditions,	
				CF3I, II, 4:1, 40 sec.,	
		erfluoro	piodopropane	(and some perfluoroiodo	opentane and a
trac	-				
	of I): I, perfluore	propene	. 5:1 by wei	ght /245°/300 atmosr	heric

of I); I, perfluoropropene, 5:1 by weight, --/245°/300 atmospheric, --, 2-trifluoromethyl-1-iodononane, bl20 129°; ICF2CF2H, II, 3:1, 3 min./260°/750 mm., 85% (13% conversion), H(CF2)4I, b750 85°; ICF2CF2Cl, II, 3:1, 3 min./260°/750 mm., 80% (9% conversion),

Cl(CF2)4I, b350 76° ; I(CF2)2I, II, 5:1, 3 min./260 $^\circ$ /750 mm., --, I(CF2)4I, b100 86° . The compds. are useful as hydraulic fluids.

L7 ANSWER 28 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1965:90276 CAPLUS

DOCUMENT NUMBER: 62:90276

ORIGINAL REFERENCE NO.: 62:16051g-h,16052a
TITLE: Perfluoroalkyl iodides

PATENT ASSIGNEE(S): E. I. du Pont de Nemours & Co.

SOURCE: 22 pp.
DOCUMENT TYPE: Patent
LANGUAGE: Unavailable

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

IF5 73.3 is mixed at room temperature with 167.7 I and 0.5 part SbF3 in a bomb jacketed with Hastelloy C. The bomb is cooled at 0°, purged of air and heated at 80° with stirring. C2F4 (I) 192 parts is added portionwise over an hr. The pressure rises from 14 to 17 kg./cm.2 after each addition After another hr. the bomb is vented through a water cleaner followed by a drying column (Drierite). The gases are collected at -60°. The non-condensed gas contains 4.3 parts C2F5I (II), while the condensed fraction (397 parts) contains 98.3% II, 0.4% I, 0.8% CF2ICF2I and 0.4% C4F10. The yield in II is 98.5% based on the consumed I. Other catalysts may be used (given: catalyst (2 parts) and yield in % II) SnF2, 99; CaF2, 78; CrF3.3H2O, 16; KF, 18; HF (100 parts), 15; CoF2.2H2O, 60; PbF4, 39; CuF2, 78; NiF2.5H2O, 28; AgF (50%), 73. Depending on the exptl. conditions, the following products were also obtained in the reaction mixture: C4F9I, C6F13I, C8F17I, C10F2II, C12F25I, C14F29I, C16F33I.

L7 ANSWER 29 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1964:410993 CAPLUS

DOCUMENT NUMBER: 61:10993

ORIGINAL REFERENCE NO.: 61:1755d-h,1756a-b

TITLE: Improvement in the preparation of

perfluoroalkyl iodides from

tetrafluoroethylene

INVENTOR(S): Parsons, Raymond E.

PATENT ASSIGNEE(S): E. I. du Pont de Nemours & Co.

SOURCE: 5 pp.
DOCUMENT TYPE: Patent
LANGUAGE: Unavailable

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 3132185		19640505	US 1962-191722	19620502
BE 640970			BE	
GB 998235			GB	

AB The preparation is given of perfluoroethyl iodide (C2F5I) (I) and longer chain title compds. C2F5(CF2CF2)nI (II), where n = 1 or greater, according to the reaction 2I2 + IF5 + 5CF2:CF2 → 5CF3CF2I. If the molar ratio of iodine to IF5 <2:1, products of formula II (mixture, various n values) result, usually in conjunction with varying amts. of I depending on the iodine to IF5 ratio. Thus, 0.66 mole iodine, 0.33 mole IF5 (III), and 0.0028 mole SbF5 were mixed at ambient temperature in a Hastelloy-C lined autoclave (2 moles iodine and 8.48 millimoles catalyst/mole III), the

whole cooled <0°, evacuated to remove air, heated to 80° with agitation and CF2CF2 (IV) and 192 parts added slowly; the pressure rose to 200-50 lb./in.2 gage in <3 min., the temperature rose $8-26^\circ$ after each addition in about 1 min. and then reverted to about 80° until 165 parts had been added. No temperature increase occurred during the addition of

remaining 27 parts of IV. The mixture was kept 1 hr. at 80°, cooled and the volatile products collected in a refrigerated receiver at -60°. Mass-sectrometry, showed 98.3% I, 0.4% unreacted IV, 0.8% IF2CCF2I (V) and 0.4% perfluorocyclobutane (VI). The noncondensed gases contained 28.1 parts IV and 4.3 parts I. The yield of I was 98.5%, based on IV. When the reaction temperature was 60° rather than 80°, the condensed product contained $9\overline{7.4}$ % I, 0.7% IV, 0.6% VI, 0.7% V, and 0.4% other products; the yield of I was 98.1%. When the reaction was repeated with omission of the SbF5, the condensed product contained 98.5% I, 0.4% IV, 0.5% V and 0.5% VI, the oily residue contained 94.6% V and 4.4% I and the yield of I was 98.4% based on IV. The reaction was repeated with 0.0128 mole anhydrous SnF2; the condensed product contained 96.9% I, 0.6% IV, 0.8% VI, 0.3% V, and 1.3% other products. The yield of I was 99% based on IV. Similarly, other catalysts were used (catalyst, and amount (parts) and % yield of I given): CaF2, 2.0, 78; CrF3.3H2O, 2.0, 16; KF, 2.0, 18; HF, 100.0, 15; CoF2.2H2O, 2.0, 60; PbF4, 2.0, 39; CuF2, 2.0, 78; NiF2.5H2O, 2.0, 28; and AgF (50%), 2.0, 73. The following examples give the mixture of components. A mixture of 0.33 mole iodine, 0.182 mole III, and 0.0028 mole SbF3 (VII) was prepared at ambient temperature in a Hastelloy-C lined autoclave,

cooled to below 0°, evacuated, sealed, heated to 60° with agitation, and 1.21 moles IV added slowly in small increments in 1 hr. The reaction was exothermic, the pressure dropped rapidly from 150-250 lb./in.2 gage to <100 lb./in.2 gage, the temperature rose 7-44° after each addition The mixture was kept at 70° 1 hr., cooled, the volatile products were vented through a H2O scrubber and a CaSO4 drying column, condensed in a refrigerated receiver at -60° and the condensed volatile material analyzed. It contained (in mole-%) 95.1 I, 1.2 C4F9I (VIII), 0.5 IV, and 1 C2F6 (IX). The noncondensed gases were found to consist of 8.0 parts IV, 4.0 parts I, and 3.1 parts IX. The nonvolatile residue (100 parts) was washed with H2O and analyzed with a vapor phase chromatograph (in weight-%): I 21.1; VIII 28.7; C6F13I (X) 20.6; C8F17I (XI) 13.0; C10F21I (XII) 7.5; C12F25I (XIII) 4.1; C14F29I (XIV) 2.3; C16F33I (XV) 1.1. The combination of these analyses indicates that 96.9% IV is accounted for; % yields are as follows (based on IV consumed): I 48.8, VIII 15.8, X 12.3, XI 1.7, XV 0.9, and IX 0.6. Thus, the total yield of I was 95.8% and the total yield of products II was 47.1%. Iodine consumption was quant. Similarly, addnl. expts. were carried out with a varying ratio of iodine and III, as well as the temperature and the catalysts.

L7 ANSWER 30 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1964:404316 CAPLUS

DOCUMENT NUMBER: 61:4316

the

ORIGINAL REFERENCE NO.: 61:684h,685a-b

TITLE: Chemistry of the metal carbonyls. XXV. Fluorocarbon

derivatives of nickel

AUTHOR(S): McBride, D. W.; Dudek, E.; Stone, F. G. A.

CORPORATE SOURCE: Harvard Univ.

SOURCE: Journal of the Chemical Society (1964), (May), 1752-9

CODEN: JCSOA9; ISSN: 0368-1769

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

AB cf. CA 60, 5532e. π -C5H5.Ni(CO)Rf [Rf = CF3, C2F5, or n-C3F7] were isolated from reactions between dicarbonyl- π -dicyclopentadienyldinickel and **perfluoroalkyl iodides**. The related complexes were also obtained, namely, π -C5H5.Ni(CO)Me, π -C5H5.Ni(Ph3P)C2F5,

and π -C5H5.Ni(Ph3P)I. Treatment of the salt K[π -C5H5.Ni(CO)] with perfluoroallyl chloride gave a mixture of π -C5H5.Ni(CO)(CF2CF:CF2) and π -C5H5.Ni(CO)(CF:CFCF3). The probable constitution of the C4F5 group in a new complex π -C5H5.Ni(Ph3P)C4F5 was discussed. Reactions between dicyclopentadienylnickel and ethylene, hexafluorobut-2-yne, and butadiene were described.

L7 ANSWER 31 OF 31 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1951:49720 CAPLUS

DOCUMENT NUMBER: 45:49720
ORIGINAL REFERENCE NO.: 45:8442a-f

TITLE: Synthesis of fluorocarbons, perfluoroalkyl

iodides, bromides, and chlorides, and

perfluoroalkyl Grignard reagents

AUTHOR(S): Haszeldine, R. N. CORPORATE SOURCE: Univ. Cambridge, UK

SOURCE: Nature (London, United Kingdom) (1951), 167, 139-40

CODEN: NATUAS; ISSN: 0028-0836

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

cf. C.A. 44, 3875h, 4861d. I(CF2CF2)2I is prepared in high yield from ICF2CF2I and C2F4 with heat, also from C2F4 and iodine by heating under pressure; in both cases higher members of the series I(CF2CF2)nI are formed in amts. that can be varied by choice of temperature and pressure. cyclic dimer of C2F4 is a by-product in these reactions. The reaction is more conveniently carried out in stages in which each member of the series is converted into the next higher member. With lower members of the series, small amts. of the compds. formed with 2 or more C2F4 mols. are produced and this tendency becomes greater with longer chains. Compds. containing up to 9 C2F4 units have been prepared A chain mechanism is proposed for these reactions. Compds. of greater chain length, from 10 to 20 C2H4 units, are obtained by heating iodine with excess C2F4 than by heating C2F4I2, C4F8I2, etc., with excess C2F4 or from C2F4 and excess iodine. Compds. with the general formula I(CF2CF2)nI have properties similar to those of CF3(CH2)nI; the lower members are liquids and the higher members are waxy solids. The compds. with n greater than 1 are much more stable than C2F4I2, but the C-I bonds are broken by heat or ultraviolet light to give the corresponding free radicals which undergo addition-polymerization with C2H4, C2H2, and yield organometallic compds. I(CF2CF2)nI with IF5, SbF5, or F diluted with N yield CF3(CF2)nI. Fluorocarbons are conveniently prepared in high yield up to C18F38 from I(CF2)nI or CF3(CF2)nI with F diluted with N, IF5, IF7, or SbF5 in an autoclave or by passing over heated CoF3. Pos. iodine is present in perfluoroalkyl mono- and diiodides and it may be replaced with H from alc. alkali or by irradiation in solvents such as hydrocarbons, alcs., and ethers. Br(CF2CF2)nBr and Cl(CF2CF2)nCl and small amts. of Br(CF2CF2)nI or Cl(CF2CF2)nI are formed by the thermal or photochem. bromination or chlorination. Perfluoroalkyl dibromides have been converted into perfluoroalkyl monobromides with BF3. Perfluoroalkyl monobromides give fluorocarbons with excess BF3 or SbF5 under pressure. The conversion of perfluoroalkyl dichlorides into perfluoroalkyl monochlorides and fluorocarbons with F diluted with N, C1F3, or SbF5 under pressure gives poorer yields than are obtained from the corresponding iodo and Br compds. Oils, greases, and waxy solids are obtained from long-chain fluorocarbons with fluorochlorocarbons or with small amts. of cyclic fluorocarbons. The lower perfluoroalkyl monoiodides form Grignard reagents that behave normally with water, CO2, and acyl halides; the latter give low yields.